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OFFICE OF CHEMICAL SAFETY AND
POLLUTION PREVENTION

MEMORANDUM

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This document provides the Health Effects Division's (HED's) risk assessment for the Expansion of Representative Commodity Use on Potato to Tuberous and Corm Vegetables Subgroup 1C.

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1.0 EXECUTIVE SUMMARY

The Interregional Research Project No. 4 (IR-4), in PP#3E8201, has proposed the expansion of the potato tolerance for residues of the herbicide paraquat (1,1'-dimethyl-4,4'-bipyridinium ion) derived from the application of the dichloride salt (calculated as the cation) to Tuberous and Corm Vegetables Subgroup 1C.

Use Profile

Paraquat dichloride (hereafter in this document referred to solely as paraquat) is a non-selective herbicide currently registered for the control of weeds and grasses in agricultural and non-agricultural areas, and for use as a defoliant, desiccant, and plant growth regulator. The Paraquat Reregistration Eligibility Decision (RED) Document (EPA 738-F-96-018) was issued in August 1997. The Product and Residue Chemistry Chapters of the RED were completed in July of 1995 (D217262, D. Miller). Currently all registered end-use products are formulated as paraquat dichloride, and the active ingredient is expressed in terms of the paraquat cation.

Tolerances for residues of paraquat have been established under 40CFR§180.205. The tolerances are expressed in terms of residues of paraquat derived from application of either the bis(methyl sulfate) or the dichloride salt (both calculated as the cation). Because there are currently no registered products containing paraquat dimethyl sulfate, the tolerance expression listed under 40CFR§180.205(a) and §180.205(c) should be revised to remove reference to the bis(methyl sulfate) salt of paraquat.

General tolerances have been established under 40CFR§180.205(a), and range from 0.01 ppm for egg and milk to 210 ppm for Animal feed, non-grass, group 18, hay. Tolerances with regional registration have been established under 40CFR§180.205(c) at 0.05 ppm for cassava, pigeon pea seed, tanager, tyfon, and true yam, and at 0.1 ppm for taro corm.

The RED Document (EPA 738-F-96-018, 8/97) recommended numerous changes to paraquat tolerances. A Federal Register notice proposing these changes was published on 8/4/2004 (69 FR 47051-47068). The proposed tolerance revisions have not as yet been finalized.

Proposed New Uses

IR-4 has submitted copies of the registered label for a 2.0 lb paraquat cation/gal emulsifiable concentrate (EC) formulation of paraquat dichloride (Gramoxone SL 2.0; EPA Reg. No. 100-1431). Gramoxone SL 2.0 is proposed for pre plant or preemergence directed spray to tuberous and corm vegetables subgroup 1C. Maximum application rate is 0.5 lb a.i./A. A maximum of three applications is proposed. No pre-harvest interval (PHI) is specified.

Hazard Identification

The toxicology database for paraquat is considered complete. The primary target organ of paraquat is the lung. Evidence of lung inflammation, scarring, and compromised lung function in response to paraquat are observed throughout the toxicity database in different species (rats, mice, and dogs). Effects in the respiratory tract are observed after acute, subchronic, and chronic exposures regardless of the route of exposure (oral or inhalation). However, inhalation was a more sensitive route of exposure than the oral route. With increasing durations of exposure,

effects of paraquat in other organ systems are observed. These effects include liver inflammation and necrosis in rats and inflammation and necrosis of the kidneys in rats and mice. Lenticular changes in the eyes of rats were also observed with increasing durations of exposure. Importantly, the lung effects occur at doses lower than effects in these other organs systems, and so protecting for lung effects protects for all other adverse effects of paraquat.

The effects of paraquat in lungs are considered systemic effects. There are no dermal toxicity studies suitable for evaluation of systemic lung effects in the toxicity database for paraquat. Therefore, the Agency is using an oral endpoint for dermal risk assessments with a dermal absorption factor of 0.3%, which was derived from dermal absorption studies conducted in humans and monkeys.

Paraquat did not cause reproductive toxicity. Developmental toxicity in response to paraquat, when observed, always occurred in the presence of maternal toxicity. Four developmental toxicity studies (two in rats and two in mice) are available. If developmental toxicity was present, clear no adverse effect levels were identified which were equivalent to (or exceeded) those for maternal toxicity. Therefore, there was no evidence of quantitative susceptibility. The kinds of developmental effects observed (e.g. reduced body weight/gain and delayed skeletal ossification) are effects that are commonly observed secondary to maternal toxicity. These developmental effects were of lesser severity than those observed in maternal animals (e.g. respiratory distress, reduced body weight, lesions in the lungs and kidneys) demonstrating no evidence of qualitative susceptibility in the young.

Previously, the Agency had required that a developmental toxicity study in rabbits be conducted for paraquat. As a result, the FQPA Safety Factor had been retained as a 3X database uncertainty factor for Females 13-39 for the acute dietary risk assessment only. The Agency reviewed the toxicity database for paraquat and concluded that a developmental toxicity study in rabbits was not likely to add information that would impact the paraquat risk assessment (TXR 0056294, April 12, 2012). Therefore, this study is no longer required and the FQPA Safety Factor has been reduced to 1X for this population.

No evidence of neurotoxicity was observed in acute and subchronic neurotoxicity studies conducted with paraquat up to the doses at which respiratory effects were observed (e.g. the maximum tolerated dose). There was also no evidence of immunotoxicity in response to paraquat.

Paraquat was found to be weakly positive in the mouse lymphoma assay and human lymphocyte cytogenetic assay, and was positive in the sister chromatid exchange assay. Conversely, paraquat was not mutagenic in the *Salmonella typhimurium* assay, was not genotoxic in the unscheduled DNA synthesis assay *in vitro* or *in vivo*, was negative for chromosomal aberration in the bone marrow test, and no evidence was found for suppressed fertility or dominant lethal mutagenicity in mice. The Cancer Peer Review Committee and the Science Advisory Committee (1989) concluded that there was no evidence of carcinogenicity in animal studies and classified paraquat as a Group E chemical (evidence of non-carcinogenicity in humans) (TXR 014110).

Paraquat is severely toxic following acute exposure via the dermal and inhalation routes (Category I) and toxic by the oral route of exposure (Category II). It is a dermal and ocular irritant but is not a skin sensitizer.

Dose Response Assessment

Toxicological points of departure (PODs) were selected for dietary/drinking water and occupational exposure scenarios for this assessment. Acute and chronic reference doses (RfDs) were selected for assessment of food and drinking water exposures. The population adjusted dose (PAD) is equivalent to the reference dose (RfD) divided by the additional FQPA Safety Factor, which was reduced to 1X for All Populations. An acute RfD/PAD for all populations was selected from a multi-generation study in rats which showed increased incidence of alveolar histiocytes in both sexes. A chronic RfD/PAD for all populations was selected from a chronic feeding study in dogs based on increased severity of chronic pneumonitis and gross lung lesions in both sexes and focal pulmonary granulomas in males. Points of departure for dermal exposures utilized a dermal absorption factor of 0.3% and the same endpoints as those utilized in the dietary assessment, with the multi-generation study in rats used for short/intermediate-term dermal assessments and the chronic feeding study in dogs used for long-term dermal assessments. A subchronic inhalation study in rats was used for short thru long-term inhalation assessments. An uncertainty factor of 100x was applied to endpoints selected for all exposure routes (10x for interspecies extrapolation, 10x for intraspecies variation).

Exposure/Risk Assessment and Risk Characterization

Risk assessments were conducted for dietary (food and water) and occupational exposure pathways based on the proposed expanded use of paraquat on Crop Subgroup 1C. Paraquat has no residential uses; therefore, a residential assessment is not required. Refined acute and chronic dietary and drinking water risk assessments for paraquat showed that dietary and drinking water exposure estimates are below HED's level of concern for the general population and all population subgroups. Occupational exposure and risk estimates indicate that worker handler and post-application exposures are not of concern at the maximum allowable application rates for the proposed expanded uses. Aggregate risks are not of concern.

Use of Human Studies

This risk assessment relies in part on data from studies in which adult human subjects were intentionally exposed to a pesticide or other chemical. These studies, listed in Appendix D, have been determined to require a review of their ethical conduct. Some of these studies are also subject to review by the Human Studies Review Board. All of the studies used have received the appropriate review. Although there are significant gaps in the ethical documentation for these studies, there is no evidence that the research was intended to harm participants or that it was fundamentally unethical in other ways (MRIDs 00126097, 00126098, & 00126099, K. Sherman, 6/11/2012).

2.0 HED RECOMMENDATIONS

2.1 Data Deficiencies/Conditions of Registration

A revised Section F is required.

2.2 Tolerance Considerations

2.2.1 Enforcement Analytical Method

PAM Vol. II lists a spectrophotometric method, designated as Method I (LOD = 0.01-0.06 ppm), as being available for the enforcement of tolerances in plant commodities. Several modifications of Method I have been developed for analysis of specific crops, and these modified methods have been used for data collection.

PAM Vol. II lists a spectrophotometric method, designated as Method Ia (LOD = 0.005 ppm), as being available for the enforcement of tolerances in animal commodities. The registrant has submitted descriptions and adequate independent laboratory validation data for a high-performance liquid chromatography method (HPLC; designated as Method 4B) to determine paraquat residues in animal tissues and eggs. The method has been validated by the Analytical Chemistry Branch (ACB). The registrant was requested to make minor changes in the method write-up. A revised version of the method has been submitted (RAM 004/04; MRID 43226902) and is now available for enforcement purposes. The reported LOQ is 0.005 ppm for livestock tissues and eggs.

2.2.2 International Harmonization

The Codex Alimentarius Commission has established maximum residue limits (MRLs) for paraquat residues in root and tuber vegetables at 0.05 ppm. The Codex and U.S. tolerances are in harmony with respect to MRL/tolerance expression; both regulate the parent paraquat cation only. The Agency cannot harmonize with the Codex MRL of 0.05 ppm since some commodities which the proposed US tolerance is translated from contained residues of > 0.05 ppm.

No Canadian or Mexican MRLs have been established for paraquat. Registered food/feed uses of paraquat exist in Canada. These uses presumably fall under the PMRA General MRL of 0.1 mg/kg. Regulation B.15.002(1) of the Canadian Food and Drugs Regulations (FDR) establishes 0.1 ppm as the General Maximum Residue Limit. This regulation states that a food is adulterated if it contains residues of a pesticide at a level greater than 0.1 ppm unless a specific MRL has been established in Table II, Division 15 of the FDR.

2.2.3 Recommended Tolerances

Pending the submission of a revised Section F, there are no residue chemistry issues that would preclude granting a registration for paraquat on the Crop Subgroup 1C, as follows:

In 2009 HED issued guidance on tolerance expressions (S. Knizner, 2009). We now conclude the tolerance expression should be as follows: *Tolerances are established for residues of paraquat, including its metabolites and degradates, in or on the commodities in the table below.*

Compliance with the tolerance levels specified below is to be determined by measuring only paraquat dichloride and calculated as the paraquat cation.

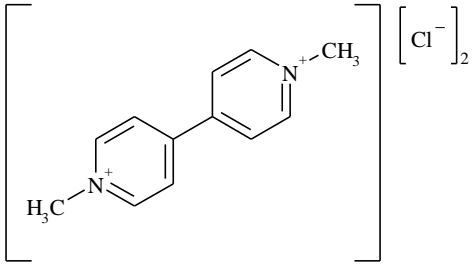
Table 1. Tolerance Summary for Paraquat Dichloride				
Commodity	Tolerance (ppm)			Comments (<i>correct commodity definition</i>)
	Established	Proposed	Recommended	
Tolerances to be established under 40 CFR ' 180.205(a):				
Tuberous and corm vegetables subgroup 1C	--	0.5	0.50	
Potato	0.5	Revoke		Covered by Tuberous and corm vegetables subgroup 1C
Tolerances to be revoked under 40 CFR ' 180.205(c):				
Cassava	0.05	Revoke		Covered by Tuberous and corm vegetables subgroup 1C
Tanier	0.05	Revoke		Covered by Tuberous and corm vegetables subgroup 1C
Yam, true, tuber	0.05	Revoke		Covered by Tuberous and corm vegetables subgroup 1C

2.2.4 Revisions to Petitioned-For Tolerances

The only revisions are for significant figures.

3.0 INGREDIENT PROFILE

3.1 Chemical Identity

Table 2. Nomenclature of Paraquat Dichloride.	
Compound	
Common name	Paraquat dichloride
IUPAC name	1,1'-dimethyl-4,4'-bipyridinium dichloride
CAS name	1,1'-dimethyl-4,4'-bipyridinium dichloride
CAS registry number	1910-42-5 (4685-14-7 for the cation)
End-use product (EP)	2.0 lb paraquat cation/gal SC (Gramoxone Inteon; EPA Reg. No. 100-1217)

3.2 Physical/Chemical Characteristics

A table of the physiochemical properties of paraquat dichloride is provided in Appendix B. Paraquat dichloride is freely soluble in water, slightly soluble in alcohols, and insoluble in nonpolar organic solvents. It has a very low vapor pressure.

3.3 Pesticide Use Pattern

3.3.1 Registered Products

There are currently 26 active paraquat dichloride registrations including 19 Section 3 registrations and six special local needs (SLNs) or 24(c) registrations and one Experimental Use Permit (EUP).

3.3.2 Proposed New Uses

IR-4 has submitted a petition (PP#3E8201) for expansion of the use on potato to Crop Subgroup 1C. The product to be used is Gramoxone SL 2.0 (EPA Reg. No. 100-1431), an emulsifiable concentrate (EC) formulation which contains 2 lb paraquat cation/gal. Gramoxone SL 2.0 is proposed for pre plant or preemergence directed spray to tuberous and corm vegetables subgroup 1C. Maximum application rate is 0.5 lb a.i./A. A maximum of three applications is proposed. No pre-harvest interval (PHI) is specified.

3.4 Anticipated Exposure Pathways

Dietary (food and water) exposures are expected based on existing uses of paraquat dichloride and the requested expanded uses. A residential exposure assessment is not required for this assessment because there are no residential uses or exposures associated with paraquat dichloride.

3.5 Considerations of Environmental Justice

Potential areas of environmental justice concerns, to the extent possible, were considered in this human health risk assessment, in accordance with U.S. Executive Order 12898, "Federal Actions to Address Environmental Justice in Minority Populations and Low-Income Populations," <http://www.eh.doe.gov/oepa/guidance/justice/eo12898.pdf>).

As a part of every pesticide risk assessment, OPP considers a large variety of consumer subgroups according to well-established procedures. In line with OPP policy, HED estimates risks to population subgroups from pesticide exposures that are based on patterns of that subgroup's food and water consumption, and activities in and around the home that involve pesticide use in a residential setting. Whenever appropriate, non-dietary exposures based on home use of pesticide products and associated risks for adult applicators and for toddlers, youths, and adults entering or playing on treated areas postapplication are evaluated. Further considerations are currently in development, as OPP has committed resources and expertise to the development of specialized software and models that consider exposure to bystanders and farm workers as well as lifestyle and traditional dietary patterns among specific subgroups.

4.0 HAZARD CHARACTERIZATION/ASSESSMENT

4.1 Toxicology Studies Available for Analysis

The toxicology database for paraquat dichloride is considered complete. Acute and subchronic neurotoxicity studies required by the new 40 CFR Part 158 data requirements have been submitted and reviewed by the Agency. An immunotoxicity study that was also a new data requirement has been submitted, been reviewed and acceptable. None of these recently submitted studies alter the endpoints selected for this risk assessment.

- Acute: Oral, dermal, inhalation, eye irritation, dermal irritation, and skin sensitization
- Subchronic: 90-day oral toxicity (dog), 21-day dermal toxicity (rabbit), 21-day inhalation
- Developmental toxicity: developmental toxicity (rat and mouse; rabbit study was waived)
- Reproduction: 3-generation reproduction (rat)
- Chronic: combined oral chronic toxicity/carcinogenicity (rat), carcinogenicity (mouse), chronic oral toxicity (dog)
- Neurotoxicity: Acute and Subchronic neurotoxicity (rat),
- Other: mutagenicity battery, metabolism, immunotoxicity (mouse), *in vivo* human dermal absorption

4.2 Absorption, Distribution, Metabolism and Excretion

Paraquat is poorly absorbed after oral administration to rats, dogs and mice. After oral administration (gastric intubation) of single doses of paraquat dichloride or dimethylsulfate to Wistar strain male and female rats, most of the administered radioactivity (69-96%) was excreted in feces as unchanged parent (e.g. was not metabolized). Of the fraction that was metabolized (up to 30%) the route of degradation was found to be microbial degradation of paraquat in the gut. Most of the administered dose of paraquat is excreted in the feces within 2-3 days. The excretion profile of paraquat changed markedly with the route of administration. After subcutaneous injection, unchanged paraquat appeared mostly in urine (73-96% of the administered radioactivity), where it was excreted as unchanged parent (73-96% dose) within 1 day of dosing.

The fraction of paraquat that is orally absorbed was rapidly distributed to most tissues (particularly the lungs and kidneys). Tissues other than the lungs did not retain paraquat.

4.3 Toxicological Effects

The primary target organ of paraquat is the lung. Evidence of lung inflammation, scarring, and compromised lung function in response to paraquat are observed throughout the toxicity database in different species (rats, mice, and dogs). Effects in the respiratory tract are observed after acute, subchronic, and chronic exposures regardless of the route of exposure (oral or inhalation). However, inhalation was a more sensitive route of exposure than the oral route. With increasing durations of exposure, effects of paraquat in other organ systems are observed. These effects include liver inflammation and necrosis in rats and inflammation and necrosis of the kidneys in rats and mice. Lenticular changes in the eyes of rats were also observed with increasing durations of exposure. Importantly, the lung effects occur at doses lower than effects in these

other organs systems, and so protecting for lung effects protects for all other adverse effects of paraquat.

The effects of paraquat in lungs are considered systemic effects. There are no dermal toxicity studies suitable for evaluation of systemic lung effects in the toxicity database for paraquat. The only available dermal toxicity study was conducted in rabbits. Since severe skin damage resulted at relatively low topical doses (6 mg/kg), it was not possible to test doses high enough to result in systemic toxicity, particularly lung effects. Therefore, the Agency is using a dermal absorption factor of 0.3% that was derived from dermal absorption studies conducted with paraquat in humans (MRID 00153439) and an oral endpoint for dermal risk assessments

Paraquat did not cause reproductive toxicity. Developmental toxicity in response to paraquat, when observed, always occurred in the presence of maternal toxicity. Four developmental toxicity studies (two in rats and two in mice) are available. If developmental toxicity was present, clear no adverse effect levels were identified which were equivalent to (or exceeded) those for maternal toxicity. Therefore, there was no evidence of quantitative susceptibility. The kinds of developmental effects observed (e.g. reduced body weight/gain and delayed skeletal ossification) are effects that are commonly observed secondary to maternal toxicity. These developmental effects were of lesser severity than those observed in maternal animals (e.g. respiratory distress, reduced body weight, lesions in the lungs and kidneys) demonstrating no evidence of qualitative susceptibility in the young.

Previously, the Agency had required that a developmental toxicity study in rabbits be conducted for paraquat. As a result, the FQPA Safety Factor had been retained as a 3X database uncertainty factor for Females 13-39 the acute dietary risk assessment only. The Agency reviewed the toxicity database for paraquat and concluded that a developmental toxicity study in rabbits was not likely to add information that would impact the paraquat risk assessment (TXR 0056294, April 12, 2012). Therefore, this study is no longer required and the FQPA Safety Factor has been reduced to 1X for this population.

No evidence of neurotoxicity was observed in acute and subchronic neurotoxicity studies conducted with paraquat up to the doses at which respiratory effects were observed (e.g. the maximum tolerated dose). There was also no evidence of immunotoxicity in response to paraquat.

Paraquat was found to be weakly positive in the mouse lymphoma assay and human lymphocyte cytogenetic assay, and was positive in the sister chromatid exchange assay. Conversely, paraquat was not mutagenic in the *Salmonella typhimurium* assay, was not genotoxic in the unscheduled DNA synthesis assay *in vitro* or *in vivo*, was negative for chromosomal aberration in the bone marrow test, and no evidence was found for suppressed fertility or dominant lethal mutagenicity in mice. The Cancer Peer Review Committee and the Science Advisory Committee (1989) concluded that there was no evidence of carcinogenicity in animal studies and classified paraquat as a Group E chemical (evidence of non-carcinogenicity in humans).

Paraquat is severely toxic following acute exposure via the dermal and inhalation routes (Category I) and is toxic by the oral route of exposure (Category II). It is a dermal and ocular irritant but is not a skin sensitizer.

The complete toxicity profile for paraquat is provided in Appendix A.

4.4 Safety Factor for Infants and Children (FQPA Safety Factor)

The toxicological database for paraquat is complete. Previously, the Agency had required that a developmental toxicity study in rabbits be conducted for paraquat. As a result, the FQPA Safety Factor had been retained as a 3X database uncertainty factor for Females 13-39 the acute dietary risk assessment. The Agency recently reviewed the toxicity database for paraquat and concluded that a developmental toxicity study in rabbits was not likely to add information that would impact the paraquat risk assessment (TXR 0056294). Therefore, this study is no longer required and the FQPA Safety Factor has been reduced to 1X for this population for the acute dietary risk assessment.

The FQPA Safety Factor remains reduced to 1X for the acute dietary assessment for the General Population and also remains reduced to 1X for the chronic dietary assessment for All Populations. Reduction of the FQPA Safety Factor to 1X is supported by:

- No evidence of neurotoxicity in the toxicity database;
- No indication of quantitative or qualitative susceptibility of mice or rats to *in utero* and/or pre- or post-natal exposure in five studies investigating these parameters;
- Clear NOAELs for developmental effects, when observed;
- A conservative dietary assessments that does not underestimate the potential exposures for infants and children;
- No registered or proposed residential uses for paraquat.

4.4.1 Completeness of the Toxicology Database

The toxicity database for paraquat is considered complete. Previously, the Agency had required that a developmental toxicity study in rabbits be conducted for paraquat in order to fully satisfy Part 158 data requirements. The Agency recently reviewed the toxicity database for paraquat and concluded that a developmental toxicity study in rabbits was not likely to add information that would impact the paraquat risk assessment (TXR 0056294). There are five studies available in which to assess the effects of paraquat on development. There are two developmental toxicity studies conducted in rats; two developmental toxicity studies conducted in mice, and a 3-generation reproductive and developmental toxicity study conducted in rats. Guideline acute and subchronic neurotoxicity studies are also available.

4.4.2 Evidence of Neurotoxicity

Guideline acute and subchronic neurotoxicity studies in adult rats were performed that included functional observational batteries for neurological effects and detailed histopathology of the nervous system. No evidence of neurotoxicity was observed in these studies at dose levels up to those that cause respiratory distress and death. Also, paraquat did not cause neurotoxic effects in any of the other studies in the toxicity database. The Agency has low concern for paraquat to be a developmental neurotoxicant due to an absence of neurotoxic effects in the Agency's toxicity database, together with developmental toxicity studies indicating that fetal toxicity is secondary to maternal toxicity. Due to this low concern, a developmental neurotoxicity study is not required.

4.4.3 Evidence of Sensitivity/Susceptibility in the Developing or Young Animal

Developmental toxicity in response to paraquat, when observed, always occurred in the presence of maternal toxicity. Four developmental toxicity studies (two in rats and two in mice) and a 3-generation reproduction toxicity study are available. Developmental toxicity was observed in one developmental rat study and one developmental mouse study and always occurred in the context of severe maternal toxicity (e.g. respiratory distress, reduced body weight, lesions in the lungs and kidneys). There was no quantitative susceptibility, since developmental effects were observed at the same dose that caused maternal toxicity. The kinds of developmental effects observed in these studies (e.g. reduced body weight/gain and delayed skeletal ossification) are effects that are commonly observed secondary to maternal toxicity, and clear NOAELs for developmental effects were always identified. Since effects in the offspring, when present, were lesser in severity than those observed in maternal animals and were also consistent with those commonly observed as secondary to maternal toxicity, the Agency has concluded that there was no evidence of qualitative susceptibility in the young to paraquat.

4.4.4 Residual Uncertainty in the Exposure Database

There are no residual uncertainties in the exposure database. The dietary risk assessment is conservative and will not underestimate dietary and/or non-dietary residential exposure to paraquat dichloride.

4.5 Toxicity Endpoint and Point of Departure

4.5.1 Dose-Response Assessment

Toxicity endpoints and points of departure (PODs) for dietary (food and water) and occupational scenarios are summarized below. A detailed description of the studies used as a basis for the selected endpoints are presented in Appendix A.

Toxicological points of departure (PODs) were selected for dietary/drinking water and occupational exposure scenarios for this assessment. Acute and chronic reference doses (RfDs) were selected for assessment of food and drinking water exposures. The population adjusted dose (PAD) is equivalent to the reference dose (RfD) divided by the additional FQPA Safety Factor, which was reduced to 1X for All Populations. An acute RfD/PAD for all populations was selected from a multi-generation study in rats which showed increased incidence of alveolar histiocytes in both sexes. A chronic RfD/PAD for all populations was selected from a chronic feeding study in dogs based on increased severity of chronic pneumonitis and gross lung lesions in both sexes and focal pulmonary granulomas in males. Points of departure for dermal exposures utilized a dermal absorption factor of 0.3% and the same endpoints as those utilized in the dietary assessment, with the multi-generation study in rats used for short/intermediate-term dermal assessments and the chronic feeding study in dogs used for long-term dermal assessments. A subchronic inhalation study in rats was used for short thru long-term inhalation assessments. An uncertainty factor of 100x was applied to endpoints selected for all exposure routes (10x for interspecies extrapolation, 10x for intraspecies variation).

4.5.2 Recommendations for Combining Exposure Routes

The acute and chronic aggregate exposure assessments for all population subgroups include only food and water exposures. There are currently no residential uses for paraquat and there are no short- or intermediate-term exposure scenarios. Therefore, non-occupational short- and intermediate-term aggregate risk assessments were not conducted.

For occupational assessments, the toxic endpoint of concern (e.g. lung effects) is the same for both inhalation and dermal routes of exposure. Therefore, these routes should be aggregated when assessing worker risks.

4.5.3 Classification of Carcinogenic Potential

Paraquat is currently placed in Category E (evidence of non-carcinogenicity to humans). The carcinogenic potential of paraquat was evaluated by the Toxicology Branch Peer Review Committee (now Carcinogenicity Assessment Review Committee (CARC)) in 1986, 1988, and 1989, and by the Scientific Advisory Panel (SAP) in 1989. In 1986 the CARC classified paraquat as a Category C carcinogen (limited evidence of carcinogenicity in animals), based on an apparent increase in erroneously combined squamous cell carcinomas in different locations in the head region. In February of 1989 the SAP classified paraquat as Category D (equivocal evidence of carcinogenicity) based on squamous cell carcinoma in the nasal cavity of 2 high-dose rats. However, the SAP also commented that endpoints other than carcinogenicity were more relevant for the regulation of paraquat. Finally, in the following month (March of 1989) the CARC placed paraquat in Category E (as it had done the previous year, 1988). As a result, for this human health risk assessment, paraquat is classified in Category E, i.e., there is evidence of non-carcinogenicity to humans (TXR 0014110).

Paraquat was found to be weakly positive in the mouse lymphoma assay and human lymphocyte cytogenetic assay and was positive for sister chromatid exchange assay. Conversely, paraquat was not mutagenic in the *Salmonella typhimurium* assay, not genotoxic in the unscheduled DNA synthesis assay *in vivo* or *in vitro*, negative for chromosomal aberration in the bone marrow test, and no evidence was found for suppressed fertility or dominant lethal mutagenicity in mice.

4.5.4 Summary of Points of Departure Used in Risk Assessment

Toxicological doses/endpoints selected for the paraquat risk assessment are provided in Table 3.

Table 3. Summary of Toxicological Doses and Endpoints for Paraquat for Use in Human Health Risk Assessments			
Exposure Scenario	Dose Used in Risk Assessment, UF	Special FQPA SF and Level of Concern for Risk Assessment	Study and Toxicological Effects
Acute Dietary (all populations)	NOAEL = 1.25 mg/kg/day UF = 100 Acute RfD = 0.0125 mg/kg/day	FQPA SF = 1x aPAD = 0.0125 mg/kg/day	Multi-generation rat study MRID 00126783, 00149748, and 00149749 LOAEL = 3.75 mg/kg/day, based on increased incidences of alveolar histiocytes in both sexes
Chronic Dietary (all populations)	NOAEL = 0.45 mg/kg/day UF = 100 Chronic RfD = 0.0045 mg/kg/day	FQPA SF = 1x cPAD = 0.0045 mg/kg/day	Chronic toxicity in dogs MRID 00132472 LOAEL = 0.93 mg/kg/day, based on increased severity of chronic pneumonitis and gross lung lesions in both sexes, and focal pulmonary granulomas in males
Dermal Short- - Intermediate - Term (1 day – 6 months)	NOAEL = 1.25 mg/kg/day	LOC = MOE = 100 Dermal absorption factor = 0.3%	Multi-generation rat study MRID 00126783, 00149748, and 00149749 LOAEL = 3.75 mg/kg/day, based on increased incidences of alveolar histiocytes in both sexes
Dermal Long-Term (> 6 months)	NOAEL = 0.45 mg/kg/day	LOC = MOE = 100 Dermal absorption factor = 0.3%	Chronic toxicity in dogs MRID 00132472 LOAEL = 0.93 mg/kg/day, based on increased severity of chronic pneumonitis and gross lung lesions in both sexes, and focal pulmonary granulomas in males
Inhalation Short - through Long - Term (1 day - > 6 months)	NOAEL = 0.01 µg/L for respirable particles NOAEL = 1.25 mg/kg/day for non-respirable particles	LOC = MOE = 100 Inhalation absorption factor = 100%	21-Day inhalation toxicity study in rats (respirable particles) MRID 00113718 LOAEL = 0.10 µg/L, based on squamous keratinizing metaplasia and hyperplasia of the epithelium of the larynx 3-generation reproduction study (non-respirable particles) MRID 00126783, 00149748, and 00149749
Cancer (oral, dermal, inhalation)	Classification: Category E (evidence of non-carcinogenicity to humans) (TXR 0014110)		

UF = uncertainty factor, FQPA SF = Special FQPA safety factor, NOAEL = no observed adverse effect level, LOAEL = lowest observed adverse effect level, PAD = population adjusted dose (a = acute, c = chronic) RfD = reference dose, MOE = margin of exposure, LOC = level of concern, NA = Not Applicable

5.0 DIETARY AND DRINKING WATER EXPOSURE AND RISK ASSESSMENT

5.1 Metabolite/Degradate Residue Profile

5.1.1 Summary of Plant Metabolism Studies

The residue chemistry chapter of the paraquat RED (D217262, D. Miller, 7/28/95) concluded that, for the purposes of reregistration and risk assessment, the qualitative nature of the residue in plants is adequately understood based on studies depicting the metabolism of paraquat in carrots and lettuce following preemergence treatment and in potatoes and soybeans following desiccant treatment. The residue of concern in plants is parent paraquat.

5.1.2 Comparison of Metabolic Pathways

Paraquat is very stable. In both primary crops and rotational crops, parent paraquat was the only major residue. In goats, pigs, and poultry, paraquat was again the only residue of concern. Paraquat was not metabolized by rats. Paraquat was poorly absorbed after oral administration to rats, dogs and mice. Once absorbed, paraquat was rapidly distributed to most tissues but especially to lungs and kidneys. Tissues other than lungs did not retain paraquat. In the environment, paraquat is very persistent and undergoes minimal degradation. As a result of the findings of the plant and animal metabolism studies as well as the environmental degradation studies, parent paraquat is the only residue of concern considered in this human health risk assessment.

5.1.3 Environmental Fate and Transport

Paraquat undergoes minimal degradation in the environment, and thus is very persistent (as parent). However, its very high propensity to bind to solids, particularly clay, makes it very immobile. In addition, paraquat does not readily appear to desorb from clay. The greatest cause for concern is likely to be erosion of contaminated sediments off-site and subsequent re-deposition onto non-target areas (especially surface water bodies). There is an additional (minor) concern for the one proposed new usage (wheat) that includes aerial spray; however, this use entails very small amounts (relative to all other uses), so spray drift onto nearby surface water drinking water sources should be fairly limited. Because of its very low mobility and strong tendency to bind tightly to soils, paraquat contamination of drinking water supplies derived from groundwater is expected to be highly unlikely. In addition, the strong binding characteristics of paraquat are likely to render most residues in raw drinking water sources removable through sedimentation processes, which are typically included as part of standard drinking water treatments.

5.1.4 Residues of Concern Summary and Rationale

The residue chemistry chapter of the RED (D217262, D. Miller, 7/28/95) concluded that for purposes of reregistration and risk assessment, the qualitative nature of the residue in plants and livestock is adequately understood based on the combined results of metabolism studies. The residue of concern is parent paraquat.

Table 5.1.4 provides a summary of the MARC decisions regarding residues of concern for paraquat.

Table 5.1.4. Summary of Metabolites and Degradates to be Included in the Risk Assessment and Tolerance Expression			
Matrix		Residues included in Risk Assessment	Residues included in Tolerance Expression
Plants	Primary Crop	Parent Paraquat	Parent Paraquat
	Rotational Crop	Parent Paraquat	Parent Paraquat
Livestock	Ruminant	Parent Paraquat	Parent Paraquat
	Poultry	Parent Paraquat	Parent Paraquat
Drinking Water		Parent Paraquat	N/A

5.2 Food Residue Profile

5.2.1 Residues in Crops

No residue field trial data have been submitted for this petition. An expansion from the potato tolerance to Crop subgroup 1C was requested. Potato is the representative crop for Crop subgroup 1C and adequate field trial data has been previously been submitted. Therefore, no additional residue data are required.

5.3 Water Residue Profile

5.3.1 Estimated Drinking Water Concentrations

The drinking water estimates used in the dietary risk assessment were provided by the Environmental Fate and Effects Division (EFED; Memo, J. Lin, 10-January-2012; D396402). EFED reviewed a non-guideline supplemental mobility study (MRID 48659501). The submitted study was conducted to evaluate the effects of traditional water treatment processes on paraquat and to determine the mobility of paraquat through soil filtration column. This memorandum only addresses the first aspect on the effects of using jar tests as a mean to mimic traditional water treatment processes to determine whether the results of jar tests are sufficient to provide the justification to refine the previous drinking water assessment (J. Lin, 11-May-2011; D381972).

¹⁴C-paraquat, spiked at ~30 ppb into the raw surface water samples from five representative US CWS (community water supply) facilities, was effectively removed by a combination of typical water treatment processes conducted on a laboratory-scale: the “laboratory jar test” (coagulation using alum with either lime or soda ash, flocculation and sedimentation), followed by dual media filtration (anthracite atop of filtering sand). The combination process was able to reduce the level of ¹⁴C-paraquat to approximate or below the limit of detection of about 0.15 µg/L (ppb). The jar test results allow EFED to better characterize potential levels in finished water for drinking water assessment purpose. The level of paraquat in the finished water of 0.15 µg/L should be used for the drinking water assessment.

5.4 Dietary and Drinking Water Exposure and Risk

Refined acute and chronic dietary and drinking water exposure and risk assessments were conducted using the Dietary Exposure Evaluation Model with the Food Commodity Intake Database (DEEM-FCID™). Dietary risk assessment incorporates both exposure and toxicity of a given pesticide. For acute and chronic dietary assessments, the risk is expressed as a percentage of a maximum acceptable dose (i.e., the dose which HED has concluded will result in no unreasonable adverse health effects). This dose is referred to as the population adjusted dose (PAD). The PAD is equivalent to the reference dose (RfD) divided by the additional Safety Factor, if applied. For acute and non-cancer chronic exposures, HED is concerned when estimated dietary risk exceeds 100% of the PAD.

5.4.1 Acute Dietary and Drinking Water Analysis

A refined (probabilistic) acute dietary exposure analysis was performed for the general population and all population subgroups. The acute analysis assumed a distribution of residues based on tolerance level residues. Empirical and DEEM default processing factors were used to modify the field trial data. Maximum screening-level percent crop treated estimates were used for commodities for which data were available. If no percent crop treated data were available, 100% crop treated was assumed. The acute analysis incorporated the jar test result concentration of 0.15 ppb for the drinking water residue. Acute dietary risk estimates are not of concern for general population or other population subgroups. The subgroup with the highest risk estimate was children 1.2 years old with a 99.9th percentile acute exposure estimate of 75% of the aPAD. The 99.9th percentile aPAD for the general U.S. population was 31%.

Table 5.4.1. Results of Acute Dietary Exposure Analysis for Paraquat (Food and Drinking Water)			
Population Subgroup	aPAD (mkd)*	99.9th Percentile	
		Exposure (mkd)	% aPAD
General U.S. Population	0.0125	0.003932	31
All Infants (< 1 year old)	0.0125	0.008424	67
Children 1-2 years old	0.0125	0.009345	75
Children 3-5 years old	0.0125	0.004594	37
Children 6-12 years old	0.0125	0.003302	26
Youth 13-19 years old	0.0125	0.002026	16
Adults 20-49 years old	0.0125	0.002108	17
Adults 50+ years old	0.0125	0.002095	17
Females 13-49 years old	0.0042	0.002060	16

*mkd: milligram per kilogram per day

5.4.2 Chronic Dietary and Drinking Water Analysis

A conservative chronic dietary exposure analysis was performed for the general U.S. population and various population subgroups. Tolerance level residues and average percent crop treated assumptions were used. DEEM default and empirical processing factors were used to modify the tolerance values. The chronic analysis incorporated the jar test result concentration of 0.15 ppb for the drinking water residue. Chronic dietary risk estimates are not of concern for general population or other population subgroups. The subgroup with the highest risk estimate were children 1-2 years old with a cPAD of 27%. The % cPAD for the general U.S. population was 7.1%.

Table 5.4.2. Results of Chronic Dietary Exposure Analysis for Paraquat (Food and Drinking Water)			
Population Subgroup	cPAD (mkd)*	Exposure (mkd)	% aPAD
General U.S. Population	0.0045	0.000321	7.1
All Infants (< 1 year old)		0.000833	19
Children 1-2 years old		0.001215	27
Children 3-5 years old		0.000819	18
Children 6-12 years old		0.000480	11
Youth 13-19 years old		0.000257	5.7
Adults 20-49 years old		0.000236	5.2
Adults 50+ years old		0.000233	5.2
Females 13-49 years old		0.000222	4.9

*mkd: milligram per kilogram per day

6.0 RESIDENTIAL EXPOSURE AND RISK ASSESSMENT

Residential exposures and risk are not assessed in this document because the proposed uses of paraquat do not involve applications by homeowners or commercial applicators in residential settings at this time and there are no existing residential uses.

6.1 Residential Bystander Postapplication Inhalation Exposure

There are no residential uses proposed for paraquat in this registration action, therefore a residential exposure assessment was not conducted.

Based on the Agency's current practices, a quantitative post-application inhalation exposure assessment was not performed for paraquat at this time primarily because of low vapor pressure ($<10^{-7}$), and the low proposed use rate (0.50 lb ai/A). However, volatilization of pesticides may be a source of post-application inhalation exposure to individuals nearby pesticide applications. The Agency sought expert advice and input on issues related to volatilization of pesticides from its Federal Insecticide, Fungicide, and Rodenticide Act Scientific Advisory Panel (SAP) in December 2009, and received the SAP's final report on March 2, 2010¹. The Agency is in the process of evaluating the SAP report and may, as appropriate, develop policies and procedures to identify the need for and, subsequently, the way to incorporate post-application inhalation exposure into the Agency's risk assessments. If new policies or procedures are developed, the Agency may revisit the need for a quantitative post-application inhalation exposure assessment for paraquat.

¹ Available: <http://www.epa.gov/scipoly/SAP/meetings/2009/120109meeting.html>

6.2 Spray Drift

Spray drift is a potential source of exposure to those nearby pesticide applications. This is particularly the case with aerial application, but, to a lesser extent, spray drift can also be a potential source of exposure from the ground application methods (e.g., groundboom) employed for paraquat. The Agency has been working with the Spray Drift Task Force (a task force composed of various registrants which was developed as a result of a Data Call-In issued by EPA), EPA Regional Offices and State Lead Agencies for pesticide regulation and other parties to develop the best spray drift management practices (see the Agency's Spray Drift website for more information).² The Agency is also taking means to qualitatively and qualitatively address spray drift as a potential source of exposure in risk assessments for pesticides through existing programs such as Ag Drift and chemical specific properties of pesticides. The potential for spray drift will be quantitatively evaluated for each pesticide during the *Registration Review* process which ensures that all uses for that pesticide will be considered concurrently.

Although a quantitative residential post-application inhalation exposure assessment was not performed as a result of pesticide drift from neighboring treated agricultural fields, an inhalation exposure assessment was performed for flaggers. This exposure scenario is representative of a worse case inhalation (drift) exposure and may be considered protective of most outdoor agricultural and commercial post-application inhalation exposure scenarios.

7.0 AGGREGATE EXPOSURE AND RISK ASSESSMENT

In accordance with the FQPA, when there are potential residential exposures to a pesticide, aggregate risk assessment must consider exposures from three major routes: oral, dermal, and inhalation. There are three sources for these types of exposures: food, drinking water, and residential uses. In an aggregate assessment, exposures from relevant sources are added together and compared to quantitative estimates of hazard (e.g., a NOAEL or PAD), or the risks themselves can be aggregated. When aggregating exposures and risks from various sources, HED considers both the route and duration of exposure. Since there are no residential uses of paraquat the acute and chronic aggregate exposures include food plus drinking water exposures. Acute and chronic aggregate risks are not of concern.

7.1 Short-, Intermediate-, and Long-Term Aggregate Risk

There are no current or proposed residential uses of paraquat. Therefore, the acute and chronic exposure estimates provided in the dietary and drinking water exposure section represent aggregate exposure.

² Available: <http://www.epa.gov/opp00001/factsheets/spraydrift.htm>

8.0 CUMULATIVE RISK

Section 408(b)(2)(D)(v) of FFDCA requires that, when considering whether to establish, modify, or revoke a tolerance, the Agency consider “available information” concerning the cumulative effects of a particular pesticide's residues and “other substances that have a common mechanism of toxicity.”

Unlike other pesticides for which EPA has followed a cumulative risk approach based on a common mechanism of toxicity, EPA has not made a common mechanism of toxicity finding as to paraquat and any other substances, and paraquat does not appear to produce a toxic metabolite produced by other substances. For the purposes of this tolerance action, therefore, EPA has not assumed that paraquat has a common mechanism of toxicity with other substances. For information regarding EPA's efforts to determine which chemicals have a common mechanism of toxicity and to evaluate the cumulative effects of such chemicals, see the policy statements released by EPA's Office of Pesticide Programs concerning common mechanism determinations and procedures for cumulating effects from substances found to have a common mechanism on EPA's website at <http://www.epa.gov/pesticides/cumulative/>.

9.0 OCCUPATIONAL EXPOSURE/RISK CHARACTERIZATION

9.1 Exposure Scenarios

Occupational handler and post-application exposure scenarios were assessed for the proposed use of paraquat on perennial tropical and sub-tropical fruit trees. Based on the product labels and information provided by the registrant, short- and intermediate-term exposure is assessed for occupational handlers and post-application activities. Dermal and inhalation exposures to workers are aggregated for paraquat because the toxicity endpoints for these exposure routes are based on common toxicological effects.

9.2 Handler Exposure

The term “handler” applies to individuals who mix, load, and apply the pesticide product. The following handler exposure scenarios were assessed for the proposed expanded uses.

9.2.1 Handler Exposure Scenarios

Based on the anticipated use patterns and current labeling, types of equipment and techniques that can potentially be used, occupational handler exposure is expected from the proposed uses. The quantitative exposure/risk assessment developed for occupational handlers is based on the following scenarios:

MIXER/LOADER

1. Open mixing/loading for groundboom and aerial applications.

APPLICATORS

1. Applying sprays with groundboom and aerial equipment.

FLAGGERS

1. Aerial applications

9.2.2 Handler Exposure Data

No chemical-specific handler exposure data were submitted in support of this registration. It is the policy of HED to use the best available data to assess handler exposure. Sources of generic handler data, used as surrogate data in the absence of chemical-specific data, include PHED 1.1, the AHETF database, the Outdoor Residential Exposure Task Force (ORETF) database, or other registrant-submitted occupational exposure studies. Some of these data are proprietary (e.g., AHETF data), and subject to the data protection provisions of FIFRA. The standard values recommended for use in predicting handler exposure that are used in this assessment, known as “unit exposures”, are outlined in the “Occupational Pesticide Handler Unit Exposure Surrogate Reference Table³”, which, along with additional information on HED policy on use of surrogate data, including descriptions of the various sources, can be found at the Agency website⁴.

9.2.3 Handler Exposure Assumptions

Unit Exposures

It is the policy of HED to use the best available data to assess handler exposure. In the case of paraquat, chemical-specific applicator exposure monitoring data were submitted to the Agency in support of the occupational handler assessment for the Paraquat Dichloride RED. For details concerning this data, see memorandums T. Manville, July 1996, D224830, D227710 and the Re-registration Eligibility Decision (RED) (August 1997). In these previous assessments, a bio-monitoring study [(MRID: 436442-02), 1995. D. Meier. Paraquat: Worker Exposure during Mixing, Loading, and Application of GRAMOXONE® EXTRA to Pecans Using Vehicle-Mounted Ground Boom Equipment] was used for assessing aerial and groundboom exposure scenarios. However, not all participants in the study adhered to label specific PPE; therefore, this study was not used in this current assessment, and instead generic handler data were used as surrogate data.

3 Available: <http://www.epa.gov/opp00001/science/handler-exposure-table.pdf>

4 Available: <http://www.epa.gov/pesticides/science/handler-exposure-data.html>

Sources of generic handler data, used as surrogate data in the absence of chemical-specific data, include PHED 1.1, the AHETF database, the Outdoor Residential Exposure Task Force (ORETF) database, or other registrant-submitted occupational exposure studies. Some of these data are proprietary (e.g., AHETF data), and subject to the data protection provisions of FIFRA. The standard values recommended for use in predicting handler exposure that are used in this assessment, known as “unit exposures”, are outlined in the “Occupational Pesticide Handler Unit Exposure Surrogate Reference Table⁵”, which, along with additional information on HED policy on use of surrogate data, including descriptions of the various sources, can be found at the Agency website⁶.

Area/Amount Treated

Based on HED ExpoSAC Policy No. 9.1, the area treated in a day was assumed to be:

- 1200 acres for mixing/loading to support aerial applications.
- 1200 acres for applying to support aerial applications.
- 200 acres for mixing/loading to support groundboom applications,
- 200 acres for applying with groundboom equipment.
- 350 acres for flaggers.

Application Rate

For the purpose of this registration action, Syngenta™ has submitted copies of the registered label for a 2.0 lb paraquat cation/gal soluble concentrate (SL) formulation of paraquat dichloride (Gramoxone 2.0; EPA Reg. No. 100-1431). Gramoxone 2.0 is proposed for post-emergence ground directed spray to previously planted crop group 1C. The maximum application rates are (2.0 pints/A) 0.50 lb a.i./A for groundboom and aerial applications, however in California, Wahsington, Oregon, and Idaho, A maximum of 3 applications is proposed along with a 14-day pre-harvest interval (PHI).

Body Weight

The average adult body weight of 80 kg was used for estimating short-term dermal and inhalation daily dose calculations.

Absorption Factors

Since the short- and intermediate-term dermal PoD was based on a 21 day inhalation study, therefore a 0.3% dermal absorption factor was used to estimate dermal exposure, and 100% absorption was assumed for estimating inhalation exposure.

Equations and Calculations

Daily Dose:

Daily dose (dermal and inhalation) is calculated by normalizing the daily exposure (dermal or inhalation) value by body weight and accounting for absorption factors:

$$\text{Average Daily Dose (mg/kg/day)} = \frac{\text{Daily Exposure (mg ai/day)} \times \text{Absorption Factor}}{\text{Body Weight (kg)}}$$

5 Available: <http://www.epa.gov/opp00001/science/handler-exposure-table.pdf>

6 Available: <http://www.epa.gov/pesticides/science/handler-exposure-data.html>

Where:

Average Daily Dose = Absorbed dose received from exposure to a pesticide in a given scenario (mg pesticide active ingredient/kg body weight/day),

Daily Exposure = Amount (mg ai/day) deposited on the surface of the skin that is available for dermal absorption or amount inhaled that is available for inhalation absorption,

Absorption Factor = A measure of the amount of chemical that crosses a biological boundary such as the skin or lungs, and

Body Weight = Body weight determined to represent the population of interest in a risk assessment.

Margin of Exposure (MOE):

The daily dermal and inhalation dose received by occupational handlers was compared to the appropriate PoD (i.e. NOAELs) to assess the risk to occupational handlers. All MOE values were calculated using the following formula:

$$MOE = \frac{NOAEL \text{ ((mg/kg/day)}}{Average \text{ Daily Dose (mg/kg/day)}}$$

Where:

MOE = Margin of exposure value used by HED to represent risk or how close a chemical exposure is to being a concern (unitless),

ADD = Average daily dose (ADD) is absorbed dose received from exposure to pesticide, and

NOAEL = Dose level in a toxicity study, where no observed adverse effects occurred in the study.

Combined Risk Estimates

Dermal and inhalation risk estimates were combined in this assessment, since the toxicological effects for these exposure routes were similar. Dermal and inhalation risk estimates were combined using the following formula:

$$Total \text{ MOE} = \frac{Point \text{ of Departure (mg/kg/day)}}{Combined \text{ dermal + inhalation dose (mg/kg/day)}}$$

9.2.4 Handler Exposure and Risk Estimates

For all short- and intermediate-term risk estimates for occupational handlers, there are no risk estimates of concern ($\text{MOE} \geq 100$). MOEs, ranged from 190 (aerial mixing and loading/baseline) to 8,800 for (groundboom “open cab” baseline with gloves). See table 9.2.4 for details.

Table 9.2.4: Occupational Short and Intermediate-Term Risk Assessment Uses of Paraquat

Table 9.2.4: Occupational Short and Intermediate-Term Risk Assessment Uses of Paraquat								
Exposure Scenario	Crops	Application. Rate	Area Treated	Inhalation Unit Exp.	Dermal Unit Exp.	Inhalation Daily Dose	Dermal Daily Dose	Combined MOE ⁵ Short/Int-term
		(lb a.i/A) ¹	(A/day) ²	(µg/lb ai) ³	(µg/lb ai) ³	(mg/kg/day) ₄	(mg/kg/day) ₄	(Dermal+ Inhalation)
Mixer/Loader (soluble concentrate)								
Aerial	Crop Group (1C)	0.50	1200	0.22 (baseline)	220 (baseline)	0.0016	0.0049	Baseline = 190
				0.0438 (label PPE ⁶)	37.6 (label PPE ⁶)	0.00033	0.00085	Baseline/Gloves/ PF5 = 1,100
Groundboom			200	0.22 (baseline)	220 (baseline)	0.00027	0.00083	Baseline = 1,100
				0.0438 (label PPE ⁶)	37.6 (label PPE ⁶)	0.000055	0.00014	Baseline/Gloves/ PF5 = 6,400
Applying (soluble concentrate)								
Groundboom (Open Cab)	Crop Group (1C)	0.50	200	0.34 (baseline)	78.6 (baseline)	0.00043	0.00030	Baseline = 1,700
				0.068 (label PPE ⁶)	16.1 (label PPE ⁶)	0.000085	0.000060	Baseline/Gloves/ PF5= 8,800
Flagger								
Aerial	Crop Group (1C)	0.50	350	0.35 (no respirator)	11 (baseline)	0.000766	0.0000724	Baseline = 1,500

1. Application rates are based on maximum values found in proposed labels: Gramoxone 2.0 @; (EPA Registration Number 100-1431)

2. Daily area treated is based on ExpoSAC Policy 9.

3. HED policies on use of surrogate data, including their sources, are presented in the “Occupational Pesticide Handler Unit Exposure Surrogate Reference Table” (<http://www.epa.gov/pesticides/science/handler-exposure-table.pdf>) and (<http://www.epa.gov/pesticides/science/handler-exposure-data.html>)

4. Daily Dose (mg/kg/day) was calculated by: [(Unit Exposure * Absorption factor * Appl. rate * Area treated) / 80 kg].

5. Short-/Intermediate-Term Combined MOE = Dermal + Inhalation. Short and Intermediate-term endpoints are the same [(NOAEL 1.25 mg/kg/day)], thus one column. The LOC for the target MOE = 100.

6. Label PPE = single layer of work clothing consisting of a long sleeved shirt, long pants, shoes, plus socks, chemical resistant gloves, protective eyewear, and a dust mist NIOSH approved respirator (N, R, P, or HE filter)

Note:

- For Aerial applicator, the groundboom applicator exposure scenario is protective.

The Agency matches quantitative occupational exposure assessment with appropriate characterization of exposure potential. While HED presents quantitative risk estimates for human flaggers where appropriate, agricultural aviation has changed dramatically over the past two decades. According the 2012 National Agricultural Aviation Association (NAAA) survey of their membership, the use of GPS for swath guidance in agricultural aviation has grown steadily from the mid 1990's. Over the same time period, the use of human flaggers for aerial pesticide applications has decreased steadily from ~15% in the late 1990's to only 1% in the most recent (2012) NAAA survey. The Agency will continue to monitor all available information sources to best assess and characterize the exposure potential for human flaggers in agricultural aerial applications.

HED has no data to assess exposures to pilots using open cockpits. The only data available is for exposure to pilots in enclosed cockpits. Therefore, risks to pilots are assessed using the engineering control (enclosed cockpits) and baseline attire (long-sleeve shirt, long pants, shoes, and socks); per the Agency's Worker Protection Standard stipulations for engineering controls, pilots are not required to wear protective gloves for the duration of the application. With this level of protection, there are no risk estimates of concern for applicators.

9.3 Post Application Exposure

HED uses the term post-application to describe exposures that occur when individuals are present in an environment that has been previously treated with a pesticide (also referred to as re-entry exposure). Such exposures may occur when workers enter previously treated areas to perform job functions, including activities related to crop production, such as scouting for pests or harvesting. Post-application exposure levels vary over time and depend on such things as the type of activity, the nature of the crop or target that was treated, the type of pesticide application, and the chemical's degradation properties. In addition, the timing of pesticide applications, relative to harvest activities, can greatly reduce the potential for post-application exposure.

For this registration action, no post-application assessment was conducted, whereas the application is a pre-emergent ground directed activity. No quantitative worker exposure is expected.

9.3.4 Restricted Entry Interval

The REI specified on the proposed label is based on the acute toxicity of paraquat. Paraquat is classified as Toxicity Category I via the dermal route and Toxicity Category I for skin irritation potential. It is not a skin sensitizer. For this registration, post-application was not assessed, whereas this action is for a pre-emergent use. The existing label (Gramazine SL 2.0) lists a 24 hr. REI for other application scenarios.

10. REFERENCES

Paraquat Dichloride. Request to Expand the Use of Paraquat Dichloride on Potato to Tuber and Corm Vegetables Subgroup 1C, T. Morton, D419828, 9/25/2014.

Paraquat Dichloride: Acute and Chronic Aggregate Dietary Exposure and Risk Assessments for the Registration Request to Expand the Use of Paraquat Dichloride on Potato to Tuber and Corm Vegetables Subgroup 1C, T. Morton, D419827, 9/25/2014.

Paraquat. Occupational and Residential Exposure Assessment for a Proposed Use on Crop subgroup 1C (tuberous and corm vegetables). J.S. Miller, D419826, 9/25/2014.

Review of Jar Test Results for Drinking Water Assessment Purpose – J. Lin, D396402, 1/10/12

APPENDICES

A TOXICOLOGY DATA SUMMARY

A.1 Guideline Data Requirements

Guideline No.	Study Type	Technical		MRID No.
		Required	Submitted	
870.3100	Subchronic (Oral) Toxicity - Rodent	N	N	00072416 00156313
870.3150	Subchronic (Oral) Toxicity - Non-Rodent	Y	Y	
870.3200	21/28-Day Dermal Toxicity	N	Y	
870.3250	90-Day Dermal Toxicity	N	N	00113718
870.3465	90-Day (21-day) Inhalation Toxicity	Y	Y	
870.3700a	Prenatal Developmental Toxicity - Rodent.....	Y	Y	00113714 43964701 00096338 43949902
870.3700b	Prenatal Developmental Toxicity - Non-Rodent.....	Y	N	
870.3800	Reproduction and Fertility Effects.....	Y	Y	
870.4100a	Chronic (Oral) Toxicity - Rodent.....	Y*	N	00132472
870.4100b	Chronic (Oral) Toxicity - Non-Rodent (Dog).....	Y	Y	
870.4200a	Carcinogenicity -Rat.....	Y*	N	00087924 40202403
870.4200b	Carcinogenicity – Mouse.....	Y	Y	
870.4300	Combined Chronic Toxicity /Carcinogenicity- Rat	Y	Y	40218001 00138637
870.6100a	Neurotoxicity - Acute Delayed Neurotox.- Hen	N	N	47794201
870.6100b	Neurotoxicity - Subchronic - Hen.....	N	N	
870.6200a	Neurotoxicity - Acute - Rat.....	Y	Y	
870.6200b	Neurotoxicity -Subchronic - Rat.....	Y	Y	47794202
870.6300	Developmental Neurotoxicity.....	N	N	48667301
870.7800	Immunotoxicity.....	Y	Y	

*Satisfied by the Combined Chronic Toxicity /Carcinogenicity- Rat Study

A.2 Toxicity Profiles

Table A2.a. Acute Toxicity Profile – Paraquat Dichloride				
Guideline No.	Study Type [species]	MRID(s)	Results ^a	Toxicity Category
870.1100	Acute oral [rat]	00054573 43685001	LD ₅₀ = 189 (M) or 125 (F) mg/kg	II
870.1200	Acute dermal [rabbit]	00054574	LD ₅₀ = 174 mg/kg (M)	I
870.1300	Acute inhalation [rat] ^b	00046105	LC ₅₀ = 1 µg/L (M/F)	I
870.2400	Acute eye irritation [rabbit]	00054575	Severe irritation	I
870.2500	Acute dermal irritation [rabbit]	00054576	Slight to severe irritation; PIS = 2.1	III
870.2600	Skin sensitization [guinea pig]	00155289	Negative	--

- ^a The test material used in the acute inhalation study was crystalline paraquat dichloride. Purity was not specified, but the purity of crystalline paraquat dichloride used in other studies was 99.9%. The test material used in the other studies was paraquat dichloride in the form of ORTHO Paraquat Concentrate 3 (end use product containing 34.4% paraquat cation). Results are expressed in terms of paraquat dichloride rather than paraquat cation.
- ^b The acute dermal (43685002), eye (43685003) and dermal (43685004) irritation and sensitization (43685005) studies are not displayed above since these studies resulted in lower toxicity and thus reduced toxicity category, likely due to a less percent of active ingredient used in the studies (D217134).

Table A2.b. Subchronic, Chronic and Other Toxicity Profile		
Guideline No./ Study Type	MRID No. (year)/ Classification/Doses	Results
870.3150 90-Day oral toxicity Beagle dog	00072416 (1981) Acceptable/guideline 0, 7, 20, 60, or 120 ppm (estimated to be 0, 0.2, 0.5, 1.5, and 3 mg/kg/day)	NOAEL = 0.5 mg/kg/day LOAEL = 1.5 mg/kg/day, based on increased lung weight and incidence of alveolitis in both sexes
870.3200 21-Day dermal toxicity New Zealand White rabbit	00156313 (1986) Acceptable/guideline 0, 0.50, 1.15, 2.60, or 6.00 mg/kg/day	Dermal NOAEL = 1.15 mg/kg/day. Dermal LOAEL = 2.60 mg/kg/day, based on small scabs at the treatment site in both sexes and epidermal erosion/ulceration, surface exudation, acanthosis, and/or inflammation in males Systemic NOAEL = 6 mg/kg/day Systemic LOAEL = not observed
870.3465 21-Day inhalation toxicity Sprague-Dawley rat	00113718 (1979) Acceptable/guideline 0, 0.012, 0.112, 0.487, and 1.280 µg/L	NOAEL = 0.012 µg/L. LOAEL = 0.112 µg/L, based on squamous keratinizing metaplasia and hyperplasia of the epithelium of the larynx.
870.3700a Prenatal developmental Wistar rat	00113714 (1978) (initial study) Acceptable/guideline 0, 1, 5, or 10 mg/kg/day	Maternal NOAEL = 1 mg/kg/day. Maternal LOAEL = 5 mg/kg/day, based on mortality, clinical signs of toxicity (piloerection, hunched posture, respiratory distress), microscopic lesions in the lungs and kidney, and decreased body weight gain (BWG). Developmental NOAEL = 1 mg/kg/day. Developmental LOAEL = 5 mg/kg/day, based on slightly decreased fetal body weights and on delayed ossification.
870.3700a Prenatal developmental Wistar rat	43964701 (1992) (subsequent study) Acceptable/guideline 0, 1, 3, or 8 mg/kg/day	Maternal NOAEL = 8 mg/kg/day (HDT). Maternal LOAEL = not observed. Developmental NOAEL = 8 mg/kg/day (HDT). Developmental LOAEL = not observed.
870.3700a Prenatal developmental SPR Alderley Park mice	00096338 (1978) (initial study) Acceptable/guideline 0, 1, 5, or 10 mg/kg/day	Maternal NOAEL = 1 mg/kg/day. Maternal LOAEL = 5 mg/kg/day based on decreased body weight gains. Developmental NOAEL = 10 mg/kg/day. Developmental LOAEL = not observed.
870.3700a Prenatal developmental CrI:CD-1 (ICR) BR mice	43949902 (1992) (subsequent study) Acceptable/guideline 0, 7.5, 15, or 25 mg/kg/day	Maternal NOAEL = 15 mg/kg/day. Maternal LOAEL = 25 mg/kg/day based on mortality, clinical signs of toxicity (piloerection, labored respiration, hunched posture, hypothermia, hypoactivity, and/or pale extremities and eyes),

Table A2.b. Subchronic, Chronic and Other Toxicity Profile		
Guideline No./ Study Type	MRID No. (year)/ Classification/Doses	Results
		<p>decreased body weights and body weight gains, increased lung weights, and gross lesions in the lung.</p> <p>Developmental NOAEL = 15 mg/kg/day. Developmental LOAEL = 25 mg/kg/day based on retardation of the skeleton and decreased fetal body weights.</p>
870.3700a Prenatal developmental – non-rodent	Waived per HASPOC memo TXR 0056294	
870.3800 Reproduction and fertility effects (3-generation) Wistar rat	00126783, 00149748, and 00149749 (1982) Acceptable/guideline 0, 25, 75, or 150 ppm (approximately equivalent to 0, 1.25, 3.75, and 7.5 mg/kg/day)	<p>NOAEL = 1.25 mg/kg/day LOAEL for parental toxicity = 3.75 mg/kg/day, based on increased incidences of alveolar histiocytes.</p> <p>Offspring NOAEL = 7.5 mg/kg/day. Offspring LOAEL = not observed.</p> <p>Reproductive NOAEL = 7.5 mg/kg/day. Reproductive LOAEL = not observed.</p>
870.4100b Chronic toxicity Beagle dog	00132472 (1983) Acceptable/guideline 0/0, 0.45/0.48, 0.93/1.00, or 1.51/1.58 mg/kg/day in males/females	<p>NOAEL = 0.45/0.48 mg/kg/day in males/females</p> <p>LOAEL = 0.93/1.00 mg/kg/day in males/females, based on increased severity of chronic pneumonitis and gross lung lesions in both sexes, and focal pulmonary granulomas in males</p>
870.4200b Carcinogenicity mouse	00087924 (1981) Acceptable/guideline 0, 0 (two controls), 12.5, 37.5, or 100/125 ppm (estimated to be 0, 0, 1.9, 5.6, and 15.0/18.8 mg/kg/day)	<p>NOAEL = 1.9 mg/kg/day.</p> <p>LOAEL = 5.6 mg/kg/day, based on decreased body weights and food consumption in females, and increased incidences of renal tubular necrosis, tubular dilatation, and interstitial nephritis in males</p> <p>No evidence of carcinogenicity</p>
870.4200b Carcinogenicity JCL:ICR mice	40202403 (1982) Acceptable/guideline 0, 2, 10, 30, or 100 ppm (estimated to be 0, 0.3, 1.5, 4.5, and 15 mg/kg/day)	<p>NOAEL = 4.5 mg/kg/day.</p> <p>LOAEL = 15 mg/kg/day, based on mortality in females</p> <p>No evidence of carcinogenicity</p>
870.4300 Chronic/Carcinogenicity Wistar rat	40218001 (1982) Acceptable/guideline 0, 6, 30, 100, or 300 ppm (equivalent to 0/0, 0.25/0.30, 1.26/1.50, 4.15/5.12, or 12.25/15.29 mg/kg/day in males/females)	<p>NOAEL = 4.15/5.12 mg/kg/day (M/F)</p> <p>LOAEL = 12.25/15.29 mg/kg/day (M/F), based on mortality</p> <p>No evidence of carcinogenicity</p>

Table A2.b. Subchronic, Chronic and Other Toxicity Profile		
Guideline No./ Study Type	MRID No. (year)/ Classification/Doses	Results
870.4300 Chronic/Carcinogenicity Fischer 344 rat	00138637, 00153223, 40202401, 40202402, and 41317401 (1983) Acceptable/guideline 0, 0 (two controls), 25, 75, or 150 ppm (estimated to be 0, 0, 1.25, 3.75, or 7.5 mg/kg/day)	NOAEL = 1.25 mg/kg/day. LOAEL =3.75 mg/kg/day, based on ocular opacity in females corroborated by lenticular changes observed microscopically No evidence of carcinogenicity
Gene Mutation 870.5100 Bacterial Gene Mutation	00100440 (1977) Unacceptable/guideline 1.0, 3.3, 10, 33, 100, 333, or 1000 µg/plate	There was no evidence of induced mutant colonies over background.
Gene Mutation 870.5100 Bacterial Gene Mutation	00100441 (1977) Acceptable/guideline 0.16, 0.8, 4, 20, 100, 500, 2500, or 5000 µg/plate	There was no evidence of induced mutant colonies over background.
Cytogenetics 870.5375 <i>In Vitro</i> Chromosome Aberration	00152692 (1985) Acceptable/guideline 0.75 to 3500 µg/mL	There was slight evidence of chromosome aberrations induced over background in the presence and absence of S9-activation.
Cytogenetics 870.5385 <i>In Vivo</i> Chromosome Aberration	40202405 (1987) Acceptable/guideline 15, 75, or 150 mg/kg (33% paraquat ion)	There was no evidence of chromosome aberration induced over background.
Other Effects 870.5550 Unscheduled DNA Synthesis	00152693 (1985) Acceptable/guideline 10 ⁻⁹ , 10 ⁻⁸ , 10 ⁻⁷ , 10 ⁻⁶ , 10 ⁻⁵ , 10 ⁻⁴ , 10 ⁻³ , or 10 ⁻² M	There was no evidence that unscheduled DNA synthesis, as determined by radioactive tracer procedures [nuclear silver grain counts] was induced.
Other Effects 870.5550 Unscheduled DNA Synthesis	40202404 (1987) Acceptable/guideline 45, 75, or 120 mg/kg (33% paraquat ion)	There was no evidence that unscheduled DNA synthesis, as determined by radioactive tracer procedures [nuclear silver grain counts] was induced.
Other Effects 870.5450 Dominant Lethal Assay	00100442 (year not reported) Acceptable/guideline 0.04, 0.4, or 4 mg/kg/day (23.8% paraquat ion)	There was no time-related positive response of increased pre- or post-implantation loss compared to controls.
Other Effects 870.5915 <i>In Vivo</i> Sister Chromatid Exchange	00152695 (1985) Acceptable/guideline 1.2, 2.5, 12.4, 24.7, 124, 247, 1240, or 2470 µg/mL	There was a concentration-related positive response of SCE induced over background in the presence of S9-activation. A positive response of SCE induced over background was also observed in the absence of S9-activation; however, there was no clear dose-response.
870.6200a Acute Neurotoxicity-rat	47794201 (2006) Acceptable/guideline 0, 25, 75, 250 mg/kg paraquat	NOAEL (neurotoxicity)= 250 mg/kg (84 mg/kg paraquat ion)

Table A2.b. Subchronic, Chronic and Other Toxicity Profile		
Guideline No./ Study Type	MRID No. (year)/ Classification/Doses	Results
	(0, 8.4, 25.1, 84 mg/kg paraquat ion) (gavage in deionized water)	
870.6200b Subchronic Neurotoxicity-rat	47794202 (2006) Acceptable/guideline 0, 15, 50, 150 ppm (0/0, 1.0/1.1, 3.4/3.9, 10,2/11.9 mg/kg/ bw/day (M/F) (mixed in diet)	NOAEL (neurotoxicity)= 150 ppm (10.2/11.9 mg/kg paraquat ion in M/F)
870.7800 Immunotoxicity [feeding]-in female B6C3F1 mice;	48667301 (2011) Acceptable/guideline 0, 25, 75, or 100 ppm (0, 6.9, 19.9, 27.3 mg/kg bw/day)	NOAEL for immunotoxicity = 100 ppm (27.3 mg paraquat dichloride/kg body weight/day). LOAEL for immunotoxicity for both AFC and NKC assays >100 ppm. Systemic NOAEL =100 ppm (27.3 mg paraquat dichloride/kg body weight/day), the highest dose tested. Systemic LOAEL for both AFC and NKC assays is not established.
Special studies Rhesus monkey and humans	00126096-00126099 (1982) Acceptable/non-guideline 607 µg intramuscular injection in monkeys or approximately 9 µg paraquat/cm ² to the skin of humans (70.0 cm ²)	Monkeys eliminated 43.5-51.5% of the administered radioactivity in the urine within 24 hours after intramuscular injection and 52.3-72.3% within 7 days post-dose. Following dermal application to humans, total urinary excretion of the applied doses was 0.052-0.702% (corrected for incomplete urinary excretion with a rhesus monkey parenteral excretion factor of 58.6%). This result suggests that the compound is poorly absorbed through the skin in humans. Peak excretion occurred during the first 24 hours post-dose.
Dermal Absorption Non-guideline	00153439 (1984)	In a dermal absorption study with healthy adult male volunteers, 0.3% of the applied ¹⁴ C paraquat dichloride was absorbed through the intact skin (forearms, and back of the hands and lower legs) during the 24-hour exposure period.

A.3 Toxicological Endpoints

A.3.1 Acute Reference Dose (aRfD) - General Population (Including Infants and Children) and Females Age 13-49

Study Selected: Reproduction and fertility effects in rats

MRID Nos.: 00126783, 00149748, and 00149749

Executive Summary: See Appendix A.4

Dose and Endpoint for Risk Assessment: NOAEL = 1.25 mg/kg/day, based on increased incidences of alveolar histiocytes in both sexes observed at the LOAEL of 3.75 mg/kg/day.

Uncertainty Factor (UF): 100x (10x for interspecies extrapolation and 10x for intraspecies variation).

Comments about Study/Endpoint: Although there was no adequate study in which a toxic endpoint could be attributed to a single acute effect, the HIARC (Memo, 4/19/2000) determined that the 3-generation reproduction study could be used for the acute RfD because the delayed toxic effects observed in this study are consistent with acute effects for paraquat poisoning in humans. This decision was confirmed by the current risk assessment team. This study should be used for all population subgroups (i.e., general population including infants and children and females age 13-49) because the dose (NOAEL = 1.25 mg/kg/day) is protective of *in utero* effects and is consistent with the maternal respiratory tract effects (edema in the alveoli and polymorphonuclear infiltration) seen in the developmental rat study at a comparable LOAEL (5.0 mg/kg/day).

$$aRfD \text{ (General Population, including infants and children)} = \frac{1.25 \text{ mg/kg (NOAEL)}}{100 \text{ (UF)}} = 0.0125 \text{ mg/kg}$$

A.3.2 Chronic Reference Dose (cRfD) - General Population (Including Infants and Children) and Females Age 13-49

Study Selected: Chronic Toxicity in Dogs

MRID No.: 00132472

Executive Summary: See Appendix A.4

Dose and Endpoint for Establishing a cRfD: NOAEL = 0.45 mg/kg/day, based on increased severity of chronic pneumonitis and gross lung lesions in both sexes, and focal pulmonary granulomas in males observed at the LOAEL of 0.93 mg/kg/day.

Uncertainty Factor(s): 100x (10x for interspecies extrapolation and 10x for intraspecies variations).

Comments about Study/Endpoint/Uncertainty Factor: This endpoint is based on the primary effect of concern (lung toxicity) and is the lowest NOAEL in the database for chronic effects via the oral route.

$$cRfD = \frac{0.45 \text{ mg/kg/day (NOAEL)}}{100 \text{ (UF)}} = 0.0045 \text{ mg/kg/day}$$

A.3.4 Incidental Oral Exposure (Short- and Intermediate-Term)

As there are no current or proposed residential uses for paraquat, the incidental oral exposure

scenario does not need to be included in this risk assessment.

A.3.5 Dermal Absorption

Dermal Absorption Factor: 0.3%

Dermal absorption was examined in a series of special studies (MRIDs 00126096-00126099). Absorption in humans was estimated to be 0.052-0.702% (corrected for incomplete urinary excretion with a rhesus monkey parenteral excretion factor of 58.6%). These studies were non-guideline and have limited use in risk assessment. However, they provide supplemental information and, considering the weight of the evidence, it was concluded that paraquat is poorly absorbed through the skin. Therefore, a 0.3% dermal absorption factor (HIARC report April 19, 2000), from a study in adult human volunteers, was selected for risk assessment. In this study, the test material was applied to the forearms as well as the backs of the hands and legs of the volunteers for 24 hours.

A.3.6 Dermal Exposure (Short-, Intermediate-, and Long-Term)

Short- and Intermediate-Term

Study Selected: Multi-generation Study in Rats

MRID Nos.: 00126783, 00149748, and 00149749

Executive Summary: See Appendix A.4

Dose and Endpoint for Risk Assessment: NOAEL = 1.25 mg/kg/day, based on increased incidences of alveolar histiocytes in both sexes observed at the LOAEL of 3.75 mg/kg/day.

Comments about Study/Endpoint: The 21-day dermal toxicity study provided a more sensitive endpoint than did the selected study. However, only localized dermal toxicity was observed in the 21-day dermal toxicity study, whereas the endpoint should be selected based on systemic toxicity. Consequently, the selected reproduction toxicity study in rats provides the most appropriate endpoint for short and intermediate exposure.

The adverse effects noted in this study were similar to those noted in the acute and short-term studies.

As an oral NOAEL was used for this endpoint, a dermal absorption factor of 0.3% (from the adult human study discussed above) should be used in risk assessment.

Dermal Exposure (Long Term)

Study Selected: Chronic Toxicity in Dogs

MRID No.: 00132472

Executive Summary: See Appendix A.4

Dose and Endpoint for Risk Assessment: NOAEL = 0.45 mg/kg/day, based on increased severity of chronic pneumonitis and gross lung lesions in both sexes, and focal pulmonary granulomas in males observed at the LOAEL of 0.93 mg/kg/day.

Comments about Study/Endpoint: The selected chronic toxicity study in dogs provides the most sensitive endpoint for long-term exposure. As an oral NOAEL was used for this endpoint, a dermal absorption factor of 0.3% should be used in risk assessment (HIARC report, April 19, 2000).

A.3.7 Inhalation Exposure (Short and Intermediate-Term)

Study Selected: 21-Day inhalation toxicity study in rats

MRID No.: 00113718

Executive Summary: See Appendix A.4

Dose and Endpoint for Risk Assessment: The NOAEL = 0.01 µg/L, based on squamous keratinizing metaplasia and hyperplasia of the epithelium of the larynx noted at the LOAEL of 0.10 µg/L.

Comments about Study/Endpoint: The lungs are a target organ of paraquat, and paraquat was most toxic when inhaled. The selected study provides the lowest toxic concentration for this time interval and exposure route. This endpoint is based on the assumption that the particle size is respirable.

A.4

EXECUTIVE SUMMARIES

90-Day Oral Toxicity – Dog

In a subchronic toxicity study (MRID 00072416), technical grade paraquat dichloride (32.2% w/w paraquat cation, Mond Reference No.: Y00061/009/004) was administered in the diet to 3 beagle dogs/sex/dose at nominal concentrations of 0, 7, 20, 60, or 120 ppm paraquat cation for up to 13 weeks. Actual intakes are estimated to be 0, 0.2, 0.5, 1.5, and 3 mg/kg/day based on Subdivision F conversion factor of 1 ppm = 0.025 mg/kg/day.

No treatment-related adverse effects were observed on ophthalmoscopic examination, hematology, clinical chemistry, or urinalysis parameters findings, or during auscultation.

At 60 ppm, absolute and relative to body lung weights were increased by 39-56% in 1 dog/sex. Alveolitis, characterized by a mixture of exudative and proliferative reactions resulting in alveolar collapse, distortion, and interstitial hypercellularity, was observed in 5/6 dogs (vs 0 controls).

The maximum tolerated dose was exceeded at 120 ppm. Two dogs/sex were sacrificed *in extremis* during the first month, suffering from marked dyspnea, harsh rales, slow and/or irregular heartbeat, and weight loss. These two dogs lost 0.90-1.20 kg. Only 1 dog/sex survived until terminal sacrifice. Decreased food consumption was noted in the female survivor. Absolute and relative to body lung weights were increased, and alveolitis was observed in all 6 dogs.

The LOAEL is 60 ppm (approximately equivalent to 1.5 mg/kg/day) based on increased lung weight and incidence of alveolitis in both sexes. The NOAEL is 20 ppm (approximately equivalent to 0.5 mg/kg/day).

This study is classified as acceptable/guideline and satisfies the guideline requirement (OPPTS 870.4100b; OECD 452) for a subchronic oral toxicity study in dogs.

870.3200 21-Day Dermal Toxicity – Rabbits

In a 21-day dermal toxicity study (MRID # not provided [Accession # 260635]), paraquat dichloride (43.5% w/w paraquat cation; Lot/Batch # SX-1465) in distilled water was applied directly to the hair-clipped intact skin of 6 New Zealand white rabbits/sex/dose at dose levels of 0, 0.50, 1.15, 2.60, or 6.00 mg/kg/day paraquat cation for 6 hours/day, 7 days/week during a 21-day period.

No treatment-related effects were observed on clinical signs, body weight, body weight gain, food consumption, on hematology or clinical chemistry parameters, or organ weights. All animals survived until scheduled sacrifice. No evidence of systemic toxicity was noted.

At 2.60 mg/kg/day, small scabs were noted at the treatment site in 2 males (Days 18 and 21) and 1 female (Days 15, 18, and 21). Microscopically evidence of dermal irritation was found in 3 males and included: epidermal erosion/ulceration, surface exudation, acanthosis, and/or inflammation.

At 6.00 mg/kg/day, very slight to well-defined erythema was noted in 4-6 rabbits/sex at Days 11, 15, 18, and 21. Small scabs were found at the treated site in 1-2 rabbits/sex on Day 11 and 12/12 rabbits at Days 15, 18, and 21. Large scabs were noted in 2-3 rabbits/sex. Grossly, crusty scabs, redness, thickened appearance, and/or prominent subcutaneous vessels were noted. Microscopically, the same lesions were observed as in the 7.8 mg/kg/day group.

The LOAEL is 2.60 mg/kg/day, based on small scabs at the treatment site in both sexes and epidermal erosion/ulceration, surface exudation, acanthosis, and/or inflammation in males. The NOAEL is 1.15 mg/kg/day.

This study is classified as acceptable/guideline and satisfies the guideline requirements (OPPTS 870.3200; OECD 410) for a 21-day dermal toxicity study in rats.

870.3465 21-Day Inhalation – Rat

In a subchronic inhalation toxicity study (MRID 00113718), Sprague-Dawley rats were exposed by whole body inhalation to paraquat dichloride (approximately 40% paraquat ion) administered as a respirable (particle sizes < 2 µm) aerosol at nominal concentrations of 0, 0.01, 0.1, 0.5, or 1.0 µg/L paraquat ion (equivalent to analytical concentrations of 0, 0.012, 0.112, 0.487, and 1.280 µg/L, respectively) for 6 hours/day, 5 days/week for 3 weeks. The numbers of rats of each sex assigned to these groups were as follows: 32 (control group); 16 (0.5 µg/L); and 36 (remaining groups). Parameters examined included clinical observations, body weights, food consumption, and water consumption. At the end of the three-week treatment period (15 total exposures), 16 rats/sex from the control group and 8 rats/sex/group from the remaining groups were terminated and examined; 8 rats/sex/group were euthanized and examined after a two-week recovery period. Gross and microscopic examinations were restricted to the respiratory tract (nasal passages, pharynx, tongue, larynx, trachea, and lungs). The remaining rats (4/sex/dose) in the control, 0.01, and 0.1 µg/L groups were euthanized after the 5th exposure, the 15th exposure, and 1, 2, and 3 days after the 15th exposure for paraquat estimations.

There were no treatment-related effects on body weights, food consumption, water consumption, or gross pathology at any concentration.

The 1.0 µg/L group was not exposed after Day 1. Of the rats in this group, 28/36 males (78%) and 29/36 females (80%) died from respiratory failure in the subsequent 14 days.

All rats in the 0.1 µg/L group exhibited nasal discharge and squamous keratinizing metaplasia, and/or hyperplasia of the epithelium of the larynx. The changes in the

epithelium were still observed in 11/16 (69%) of the rats euthanized at the end of the recovery period.

Additionally in the 0.5 µg/L group, the following findings were observed after 3 weeks: (i) extensive ulceration, necrosis, inflammation and squamous keratinizing metaplasia, and marked/moderate hyperplasia of adjacent epithelia in larynx of all rats; and (ii) aggregations of foamy macrophages in the bronchioles or alveoli, hypertrophy of the epithelium and thickened alveolar walls in the lungs of most or all rats. After the 2-week recovery period, no ulceration or necrosis was observed in the larynx, but changes in the lungs were still seen. In addition, disruption of bronchiolar epithelium, adjacent to the macrophage aggregation, was noted.

At 0.01 µg/L, there were no treatment-related effects on any parameter.

The LOAEL is 0.10 µg/L based on squamous keratinizing metaplasia and hyperplasia of the epithelium of the larynx. The NOAEL is 0.01 µg/L.

At the request of the Agency, this study was conducted for a duration of three weeks, instead of the 90 days required by Guideline OPPTS 870.3465. Aside from the different study duration, this study was conducted in accordance with Guideline OPPTS 870.3465.

This 21-day inhalation toxicity study is classified as acceptable/guideline and satisfies the guideline requirement (OPPTS 870.3465; OECD 413) for a subchronic inhalation study in the rat.

Prenatal Developmental Toxicity

870.3700a Prenatal Developmental Toxicity Study – Rat

In a developmental toxicity study (MRID 00113714), paraquat dichloride (100% technical grade; Batch # ADYM76/C; 38% w/v paraquat ion) in 0.5% aqueous Tween 80 was administered daily via oral gavage to 29-30 presumed pregnant Alderly Park Wistar-derived (Alpk:SPF SD) rats/group at a dose volume of 10 mL/kg at dose levels of 0, 1, 5, or 10 mg/kg/day of paraquat ion from gestation day (GD) 6 through 15. All surviving dams were killed on GD 21. The lungs and kidneys from at least 11 surviving dams/group were examined microscopically. The fetuses were removed by cesarean section and examined.

One 5 mg/kg/day dam had excessive blood loss from the vagina and was euthanized on GD 18. At ≥ 5 mg/kg/day, clinical signs of toxicity included piloerection, weight loss, hunched appearance, and respiratory distress. Earliest onset of these signs occurred on GD 8, with weight loss, thin appearance, and slightly hunched posture observed in a single animal (#71); and slight piloerection was noted in another dam (#83) on the following day. Clinical signs of toxicity became more prevalent, affecting more animals more frequently, from GD 13-21. Body weight gains were decreased by 37-74% during the treatment (GD 6-16) interval (calculated by the reviewers; statistics not performed)

and by 24-29% for the overall (GD 0-21) study ($p \leq 0.001$).

Additionally at 10 mg/kg/day, one female (# 92) delivered prematurely on GD 21. An additional 14 dams died or were sacrificed moribund prior to scheduled termination. Morbidity considered to be due to the test substance was observed in 6 of these dams. Gross necropsy indicated that the lungs were red and patchy, and microscopic examination revealed large amount of edema fluid and polymorph infiltration in the alveoli, while the kidneys showed widespread degenerative change in the proximal tubules. It was stated that the deaths of the remaining dams in this dose group were likely due to gavage error.

The maternal LOAEL is 5 mg/kg/day based on mortality, clinical signs of toxicity, and decreased body weight gains. The maternal NOAEL is 1 mg/kg/day.

There was no effect on the proportion of dams having one or more resorptions, and there were no treatment-related effects on sex ratio or embryonic or fetal survival. There were no increases in fetal external visceral, or skeletal malformations or variations at any dose tested, indicating that paraquat dichloride is not teratogenic in rats at the dose levels tested.

At ≥ 5 mg/kg/day, fetal body weights were reduced by 3-6%. Skeletal ossification was slightly retarded in these groups, as indicated by decreased ossification of the caudal vertebrae and decreased degree of ossification in the digits in the fore- and hind-limbs. The percent of fetuses with 7 or 8 caudal vertebrae ossified was decreased ($p \leq 0.05$) at this dose (8% treated vs 26% controls). The percent of fetuses with "good" (Grade 2) ossification in the digits in the fore-limbs was dose-dependently decreased at 5 (29%) and 10 (23%) mg/kg/day compared to controls (42%). The percent of fetuses with Grade 2 or 3 ossification in the digits in the hind-limbs was dose-dependently decreased at ≥ 5 mg/kg/day (20% each treated) compared to controls (42%). Likewise, the percent of fetuses with "poor" (Grade 5) ossification in the digits of the hind limbs was increased at ≥ 5 mg/kg/day (23-32%) compared to controls (13%). These decreases in growth and development are probably associated with the maternal toxicity observed at this dose.

The developmental LOAEL is 5 mg/kg/day based on slightly decreased fetal body weights and on delayed ossification. The developmental NOAEL is 1 mg/kg/day. This study is classified acceptable/guideline and satisfies the guideline requirements (OPPTS 870.3700a) for a developmental study in the rat.

In a developmental toxicity study (MRID 43964701), paraquat dichloride (38.2% w/v paraquat ion) in deionized water was administered daily via oral gavage to 24 presumed pregnant Alderley Park, Wistar-derived (Alpk:APfSD) rats/group at a dose volume of 10 mL/kg at dose levels of 0, 1, 3, or 8 mg/kg/day from gestation day (GD) 7 through 16. All surviving dams were killed on GD 22. The fetuses were removed by cesarean section and examined.

There were no effects of treatment on mortality, clinical signs, body weights, body

weight gains, food consumption, or gross pathology.

The maternal LOAEL was not observed. The maternal NOAEL is 8 mg/kg/day (highest dose tested).

There were no premature deliveries or complete litter resorptions and no effects of treatment on the numbers of live fetuses, early resorptions, late resorptions, or post-implantation loss, indicating no effect on embryonic or fetal survival. In the fetuses, there were no treatment-related external, visceral, or skeletal malformations or variations, indicating that paraquat dichloride is not teratogenic in rats at the dose levels tested.

The developmental LOAEL was not observed. The developmental NOAEL is 8 mg/kg/day (highest dose tested).

A LOAEL was not observed in this study, but the dose range tested did not include the limit dose (1000 mg/kg/day). However, maternal rats exhibited mortality, clinical signs of toxicity, and microscopic lesions in the lungs and kidney in a previous study (MRID 00113714) conducted by the performing laboratory in 1978 using the same strain of rat. Because death was observed at 5 and 10 mg/kg/day in the previous study, the dose levels for the current study were lowered slightly to a maximum dose level of 8 mg/kg/day. Thus, the current study is acceptable for regulatory purposes. For the purpose of risk characterization, the NOAEL from the older study should continue to be used until additional data have been received to support any changes.

This study is classified acceptable/guideline and satisfies the guideline requirements (OPPTS 870.3700a) for a developmental study in the rat.

870.3700b Prenatal Developmental Toxicity Study – Mouse

In a developmental toxicity study (MRID 00096338), paraquat dichloride (100% technical grade; 38% w/v paraquat ion) in 0.5% aqueous Tween 80 was administered daily via oral gavage to 30 mated female SPF Alderley Park mice/group at a dose volume of 10 mL/kg at dose levels of 0, 1, 5, or 10 mg/kg/day of paraquat ion from gestation day (GD) 6 through 15. Because several mice in each group either died, littered early, or were not pregnant, an insufficient number of litters were available for teratology assessment. Therefore, 4-5 weeks after the first matings, an additional 42 mice were mated; and 6, 6, 20, and 10 mice were allocated to the 0, 1, 5, or 10 mg/kg/day groups, respectively. All surviving dams were killed on GD 18. The lungs and kidneys from at least 8 surviving dams/group were examined microscopically. The fetuses were removed by cesarean section and examined.

There were no treatment-related effects on mortality, clinical signs, food consumption, water consumption, gross pathology, or histopathology. Body weight gains for the treatment period (GD 6-15) were decreased by 22% at 5 mg/kg/day and by 13% at 10 mg/kg/day, resulting in decreased body weight gains for the overall (GD 0-18) study of 14% ($p \leq 0.05$) at 5 mg/kg/day and 11% (not significant) at 10 mg/kg/day.

The maternal LOAEL is 5 mg/kg/day based on decreased body weight gains. The maternal NOAEL is 1 mg/kg/day.

The numbers of viable fetuses and resorptions and the sex ratio, fetal weights, and litter weights in the treated groups were comparable to controls. There were no treatment-related fetal external, visceral, or skeletal malformations, variations, or retardations.

The developmental LOAEL was not observed. The developmental NOAEL is 10 mg/kg/day.

This study is classified acceptable/guideline and satisfies the guideline requirements (OPPTS 870.3700a) for a developmental toxicity study in the mouse.

In a developmental toxicity study (MRID 43949902), paraquat dichloride (38.2% w/v paraquat ion) in deionized water was administered daily via oral gavage to 26 mated female Crl:CD-1 (ICR) BR mice/group at a dose volume of 10 mL/kg at dose levels of 0, 7.5, 15, or 25 mg paraquat ion/kg/day from gestation day (GD) 6 through 15. All surviving dams were killed on GD 18. The lungs (with trachea) and kidneys were removed, weighed, and fixed in buffered formal saline, but were not examined microscopically. The fetuses were removed by cesarean section and examined.

There were no treatment-related effects on maternal food consumption.

At 25 mg/kg/day, one dam was found dead on GD 16, with no clinical signs of toxicity observed prior to death. Four other dams at this dose were euthanized prior to scheduled termination due to poor condition on GD 15-17. Clinical signs of toxicity in these dams included piloerection, labored respiration, hunched posture, hypothermia, hypoactivity, and/or pale extremities and eyes. Additionally at this dose, body weights were decreased by 6-9% ($p<0.05$) during GD 15-18 compared to controls. Body weight gains were decreased ($p<0.01$) by 22% for the treatment period (GD 6-15) and by 29% for the post-treatment period, resulting in decreased ($p<0.01$) body weight gains for the overall (GD 0-18) study of 19%. When adjusted for gravid uterine weight, body weight gains were still decreased by 18% (not significant). Dark red lung lobes were observed in the dam that was found dead, in all four of the dams sacrificed in moribund condition, and in four additional dams at termination (for a total of 35% treated vs 0% controls). Absolute and relative (to body) lung weights were increased ($p<0.01$) by 31-64% in these animals.

There were no effects of treatment at 7.5 or 15 mg/kg/day.

The maternal LOAEL is 25 mg/kg/day based on mortality, clinical signs of toxicity, decreased body weights and body weight gains, increased lung weights, and gross lesions in the lung. The maternal NOAEL is 15 mg/kg/day.

There were no premature deliveries or complete litter resorptions and no effects of treatment on the numbers of litters, fetuses (live or dead), resorptions (early or late), or on

sex ratio or post-implantation loss. There were no treatment-related fetal external, visceral, or skeletal malformations.

At 25 mg/kg/day, fetal body weights were decreased ($p < 0.01$) by 9-10% compared to controls. Retardation of the skeleton was indicated by increases ($p < 0.05$) in the numbers of: litters with retarded ossification of the occipital bone (42.9% treated vs 8.3% controls); fetuses and litters with ≤ 6 caudal centra (47.0% fetuses in 57.1% treated litters vs 7.0% fetuses in 20.8% control litters); litters with uni- or bi-lateral extra 14th ribs (64.3% treated vs 29.2% controls); and fetuses and litters with non-ossified astragalus in the hind limb (36.1% fetuses in 57.1% treated litters vs 8.9% fetuses in 20.8% control litters).

There were no effects of treatment at 7.5 or 15 mg/kg/day.

The developmental LOAEL is 25 mg/kg/day based on retardation of the skeleton and decreased fetal body weights. The developmental NOAEL is 15 mg/kg/day.

This study is classified acceptable/guideline and satisfies the guideline requirements (OPPTS 870.3700a) for a developmental toxicity study in the mouse.

Reproductive Toxicity

870.3800 Reproduction and Fertility Effects – Rat

In a three-generation reproduction toxicity study (MRIDs 00126783, 00149748, and 00149749), technical grade paraquat dichloride (32.7% w/v paraquat cation) was administered continuously in the diet to Wistar-derived Alderley Park rats (15 males and 30 females/dose) at dose levels of 0, 25, 75, or 150 ppm (approximately equivalent to 0, 1.25, 3.75, and 7.5 mg paraquat ion/kg/day, assuming that for an older rat, 1 ppm = 0.05 mg/kg/day). Parents were fed test diets for 11-12 weeks before they were mated to produce the F1, F2, and F3 litters. The F1a pups were weaned on postnatal day (PND) 21, but were not bred. P generation rats were re-mated 7 days later to produce the F1b litters, which were weaned on PND 28 and housed until PND 35. From the F1b litters, 15 males and 30 females/dose were fed the test diets and bred for the production of the F2a and F2b litters. This process was repeated to produce the F3a and F3b litters. The study was terminated after the F3b litters were weaned.

There were no effects of treatment on clinical signs, body weights, body weight gains, food consumption, food utilization, ophthalmology, hematology, clinical chemistry, or urinalysis.

In all generations, alveolar histiocytosis was increased in the 75 (10-71%) and 150 (50-86%) ppm males compared to controls (11-30%) and in the 75 (62-80%) and 150 (80-100%) ppm females compared to controls (28-40%).

High mortality was observed in the 150 ppm P, F1, and F2 females (17-43%) compared

to controls (0-4%). These deaths were considered to be due to severe lung damage caused by paraquat. The incidence of lung lesions (red or purple discoloration, congestion, edema, fibrosis, hyaline membrane formation, inflammatory cell infiltration, and/or hyperplasia) ranged from 27-35% in these animals compared to 0 controls.

At termination, the most frequent microscopic findings were hydronephrosis, nephrocalcinosis, lung congestion and/or alveolar hemorrhage, perivascular inflammatory cell infiltration in the lungs, focal accumulation of lymphocytes in the liver, and hypoplasia, atrophy, and/or necrosis of the testes. However, the incidences of these findings were not dose-related.

The LOAEL for parental toxicity is 75 ppm (approximately equivalent to 3.75 mg paraquat ion/kg/day), based on increased incidences of alveolar histiocytes in both sexes. The NOAEL is 25 ppm (equivalent to 1.25 mg paraquat ion/kg/day).

There were no effects of treatment on maternal neglect index (% dams with all pups dead by PND 10), lactation index (i.e., survival to PND 21), litter size (viability) from PND 0 through 28, or litter weight gain.

The LOAEL for offspring toxicity was not observed. The NOAEL is 150 ppm (approximately equivalent to 7.5 mg/kg/day).

There were no effects of treatment on fertility, gestation duration, or live birth index.

The LOAEL for reproductive toxicity was not observed. The NOAEL is 150 ppm (approximately equivalent to 7.5 mg/kg/day).

This study is classified as acceptable/guideline and satisfies the guideline requirements (OPPTS 870.3800; OECD 416) for a multi-generation reproduction study in the rat.

Chronic Toxicity

870.4100a (870.4300) Chronic Toxicity – Rat

In this combined chronic toxicity/carcinogenicity study (MRIDs 00138637, 00153223, 40202402, 40202401, and 41317401), paraquat dichloride (96.1% a.i.; Batch #: S 358) was administered in the diet to 70 Fischer 344 rats/sex/dose at nominal concentrations of 0, 0 (two controls), 25, 75, or 150 ppm for up to 117 weeks in males and 124 weeks in females. All doses are for the paraquat cation, and group mean actual intakes for the entire study period were not reported. Actual intakes are estimated by the reviewers to be 0, 0, 1.25, 3.75, and 7.5 mg/kg/day based on Subdivision F conversion factor of 1 ppm = 0.050 mg/kg/day. Ten rats/sex/dose were sacrificed at Week 52, and paraquat levels were determined in the tissues and plasma of 5 rats/sex/dose.

No adverse, treatment-related effects were observed on mortality, food consumption, water consumption, or on any hematological, clinical chemistry, or urinalysis parameters.

At 150 ppm, body weights were decreased ($p \leq 0.05$) at Weeks 26, 52, 78, 104, and 113 by 6-8% in males and at Weeks 52, 78, 104, 117, and 122 by 2-10% in females. Body weight gain (Weeks 0-104) were decreased by 11% in males and 9% in females (calculated by reviewers. Food utilization was decreased during Weeks 13-26, 27-40, and 41-52 (last week calculated) by 8-21% in both sexes.

The eyes were a target organ. During clinical observations, eye opacity was observed in the 75 and 150 ppm females (23-58% treated vs 5% controls) and 150 ppm males (37% vs 7%). Opacities were first observed during the first year, but were rare until after Week 101 in males and Week 111 in females. During ophthalmoscopic examinations, increased incidences (% treated vs % controls) of the following findings were observed at 150 ppm: (i) posterior polar opacity/cataract in males (41% vs 7%) and females (64% vs 0%); (ii) posterior capsular opacity/cataract in males (52% vs 0%) and females (26% vs 4%); (iii) radial cataract in both sexes (11-17% vs 0%); (iv) cataracts in both sexes (9-11% vs 2%); and (v) lens resorption in both sexes (6-9% vs 0%). Increased incidences of lens lesions were also observed in the 150 ppm group at Weeks 110 and at termination. Only minor increases in eye lesions were noted before Week 103. Microscopically, increased incidences of the following ocular lesions were observed in all treated groups (both sexes) at termination of the study: peripheral Morganian corpuscles, peripheral lenticular degeneration, mid-zonal lenticular degeneration, and a pear-shaped posterior peripheral lenticular change. Clinical observations and ophthalmoscopic examinations did not corroborate ocular toxicity at 25 ppm. Additionally, increased incidences of the following ocular lesions were noted in the 150 ppm males: lens capsule fibrosis, lens capsule rupture, peripheral retinal degeneration of the outer nuclear layer, posterior synechia, proteinaceous aqueous humor, and vitreous cellularity.

At 150 ppm, the lungs were also a target organ. At the interim sacrifice, an increase in alveolar pigmented macrophages was observed in the females (40% treated vs 20% controls). Relative to body lung weights were increased by 14-16% in both sexes at study termination. Grossly, increased incidences of occasional or multiple, dark or pale subpleural foci/areas were observed at the terminal sacrifice in males (21/33 treated vs 2/60 controls) and females (22/29 treated vs 10/58 controls). Increased incidences ($n=55-60$) were noted of alveolar epithelialization (16% vs 3%) and increased number of macrophages (17% vs 2%) in males and accumulation of alveolar macrophages in females (37% vs 19%).

The following observations were considered equivocal due to a lack of corroborating evidence of toxicity: an increased incidence ($n=55-60$) of degeneration of sciatic nerve fibers was observed in the 75 and 150 ppm males (53-54% treated vs 32% controls); an increased incidence of hydrocephalus was observed in the 75 and 150 ppm females (21-34% treated vs 12% controls).

Small amounts of the paraquat cation were detected in one or more treated groups in the lungs, liver, kidneys, skin, and plasma in animals sacrificed after 52 weeks of treatment. At the high dose, concentrations ranged from 0.037-0.71 $\mu\text{g/g}$.

The LOAEL is 75 ppm (approximately equivalent to 3.75 mg/kg/day), based on ocular opacity in females corroborated by lenticular changes observed microscopically. The NOAEL is 25 ppm (approximately equivalent to 1.25 mg/kg/day).

At the doses tested, there was a treatment related increase in tumor incidence when compared to controls. The incidences of the following tumors were increased (n=69-70; % in treated group[s] vs concurrent controls vs reference range of the animal provider): pulmonary adenomas in the 75 and 150 ppm males (6-7% treated vs 1% controls vs 1.4-5.6% reference) and in the 75 and 150 ppm females (3-11% treated vs 0% controls vs 0.8-1.7% reference); pulmonary carcinoma in the 150 ppm males (4% treated vs 1% controls vs 0.8-2.2% reference) and all treated female groups (1-3% treated vs 0% controls vs 0% reference). Additionally in the 150 ppm males at the terminal sacrifice (n=23-33), increased incidences were noted of benign pheochromocytoma in the adrenals (27% treated vs 10% controls) and thyroid parafollicular adenoma (33% treated vs 17% controls). Dosing was considered adequate based on eye and lung toxicity, and reduced body weights, body weight gains, and food utilization.

This study is acceptable/guideline and satisfies the guideline requirement for a chronic/carcinogenicity study (OPPTS 870.4300; OECD 453) in rats.

870.4100b Chronic Toxicity – Dog

In a chronic toxicity study (MRID 00132472), technical grade paraquat dichloride (32.3% w/w paraquat cation, Mond Reference No.: S358/2) was administered in the diet to 6 beagle dogs/sex/dose at nominal concentrations of 0, 15, 30, or 50 ppm (equivalent to 0/0, 0.45/0.48, 0.93/1.00, and 1.51/1.58 mg/kg/day paraquat cation in males/females) for up to 52 weeks.

No treatment-related adverse effects were observed on mortality, body weights, body weight gains, or on ophthalmoscopic examination, hematology, clinical chemistry or urinalysis parameters.

Increased incidences of the following clinical signs were observed at 50 ppm in both sexes: hypernea (4/6 vs 1/6, each sex), increased vesicular sound (3-4/6 vs 0/6), and reddening of tongues (6/6 vs 4/6, each sex). The frequency of these observations was also increased at 50 ppm. These signs were first observed at Week 13 (hypernea and increased vesicular sound) and week 9 (tongue reddening). Food consumption was decreased in one 50 ppm dog/sex. The hypernea was corroborated by further findings of pulmonary toxicity. The other findings are considered equivocal.

Lungs were the target organ. Absolute and relative to body lung weight were each increased by 36% in males and 61% in females at 50 ppm. Chronic pneumonitis was observed in 44 of the 48 dogs that were evaluated; therefore, an increased incidence was not observed. However, an increase in severity was observed in the 30 and 50 ppm

groups; the incidence (# affected/6, treated vs controls) of slight to marked chronic pneumonitis was 5-6 treated males vs 2 controls and 3-6 treated females vs 1 control. This lesion correlated to yellow discoloration and consolidation of areas of the lungs observed grossly. Additionally, the incidence and severity of minimal to moderate focal granuloma was increased in the 30 and 50 ppm males (5/6 each treated vs 4/6 controls). Focal pleural fibrosis was observed in 3/6 males at 50 ppm vs 2/6 controls and may have been treatment-related.

Small amounts of the paraquat cation were detected in the lungs of all treated groups (0.13-1.04 µg/g) and in the kidney of the 30 and 50 ppm groups (0.12-0.19 µg/g).

The LOAEL is 30 ppm (equivalent to 0.93/1.00 mg/kg/day in males/females) based on increased severity of chronic pneumonitis and gross lung lesions in both sexes, and focal pulmonary granulomas in males. The NOAEL is 15 ppm (equivalent to 0.45/0.48 mg/kg/day in males/females).

At the doses tested, there was no treatment-related increase in tumor incidence when compared to controls. Dosing was considered adequate based on an increase in pulmonary toxicity.

This study is classified as acceptable/guideline and satisfies the guideline requirement (OPPTS 870.4100b; OECD 452) for a chronic oral toxicity study in dogs.

Carcinogenicity

870.4200a Carcinogenicity Study – rat

In this combined chronic toxicity/carcinogenicity study (MRID 40218001), paraquat dichloride (≥98% a.i.; Lot #: 540108) was administered in the diet to 50 Wistar rats/sex/dose at nominal concentrations of 0, 6, 30, 100, or 300 ppm (equivalent to 0/0, 0.25/0.30, 1.26/1.50, 4.15/5.12, and 12.25/15.29 mg/kg/day in males/females) for up to 104 weeks. Additionally 12 rats/sex/dose were treated similarly, and 6 rats/sex/dose were sacrificed at Weeks 26 and 52.

No adverse, treatment-related effects were observed on body weight, body weight gains, food consumption, or on any ophthalmoscopic examination, hematological, clinical chemistry, or urinalysis parameters, organ weights, or gross and histological pathology.

Increased mortality was observed in the 300 ppm males (↑26%) and females (↑10%). In moribund animals, decreased spontaneous mobility, loss of coat luster, and piloerection were noted.

The LOAEL is 300 ppm (equivalent to 12.25/15.29 mg/kg/day), based on mortality. The NOAEL is 100 ppm (approximately equivalent to 4.15/5.12 mg/kg/day).

At the doses tested, there was no treatment-related increase in tumor incidence when compared to controls. Dosing was considered adequate based on decreases in survival in both sexes.

This study is acceptable/guideline and satisfies the guideline requirement for a chronic/carcinogenicity study (OPPTS 870.4300; OECD 453) in rats.

870.4200b Carcinogenicity (feeding) – Mouse

In a carcinogenicity study (MRID 00087924), 60 Swiss-derived mice/sex/dose were exposed to paraquat dichloride (96.1% a.i.; Batch #: S 358/1) in the diet at nominal concentrations of 0, 0 (two controls), 12.5, 37.5, or 100/125 ppm for up to 99 weeks. All doses are for paraquat cation, and actual intakes were not reported. Actual intakes were estimated to be 0, 0, 1.9, 5.6, and 15.0/18.8 mg/kg/day based on Subdivision F conversion factor of 1 ppm = 0.150 mg/kg/day. Animals in the high dose group received 100 ppm for the first 35 weeks and 125 ppm for the remainder of the study because no signs of toxicity were observed at 100 ppm. An additional 10 mice/sex/dose were treated similarly, and paraquat levels were determined in the tissues and plasma at 52 weeks.

Blood and urine analysis (except for determination of paraquat levels) were not performed and no organ was weighed. No treatment-related effect was observed on food utilization.

At 37.5 ppm, the following findings were noted: in females, decreased body weights beginning at Week 68 (↓5-20%) and decreased food consumption at Weeks 6-20 (↓3-15%) and 56-84 (↓15-22%); in males, increased incidences (% treated vs % controls, observed at terminal kill) of renal tubular necrosis (38% vs 8%), tubular dilatation (8% vs 0%), and interstitial nephritis (23% vs 15%).

At 100/125 ppm, the findings at 37.5 ppm were more severe, and the following additional findings were observed: increased incidences of hypercellularity of alveolar walls, renal tubular necrosis, tubular dilatation, and pelvic dilatation in both sexes; lung congestion and alveolar macrophages in females; and also increased mortality in females.

Small amounts of the paraquat ion were detected at 100/125 ppm in the plasma (0.051-0.056 µg/ml), kidneys (1.17-1.61 µg/g), and lungs (0.43-0.52 µg/g) of both sexes. Paraquat cation levels in other tissues were not reported.

The LOAEL is 37.5 ppm (approximately equivalent to 5.6 mg/kg/day) based on decreased body weights and food consumption in females, and increased incidences of renal tubular necrosis, tubular dilatation, and interstitial nephritis in males. The NOAEL is 12.5 ppm (approximately equivalent to 1.9 mg/kg/day).

At the doses tested, there was no treatment-related increase in tumor incidence when compared to controls. Dosing was considered adequate based on decreases in survival,

body weights, food consumption, nephrotoxicity, and lung toxicity.

This study is classified as acceptable/guideline and satisfies the guideline requirements (OPPTS 870.4200b; OECD 451) for a carcinogenicity study in mice.

In a carcinogenicity study (MRID 40202403), 80 JCL:ICR mice/sex/dose were exposed to paraquat dichloride ($\geq 98\%$ a.i.; Lot #: 540108) in the diet at nominal concentrations of 0, 2, 10, 30, or 100 ppm for up to 104 weeks. All doses are for paraquat cation, and actual intakes were not reported. Actual intakes were estimated to be 0, 0.3, 1.5, 4.5, and 15 mg/kg/day based on Subdivision F conversion factor of 1 ppm = 0.150 mg/kg/day. Interim sacrifices of 10 mice/sex/dose were performed at 26 and 52 weeks.

No treatment-related effect was observed on body weights, body weight gains, food consumption, food efficiency, or on hematology, clinical chemistry, or urinalysis findings, organ weights, or gross or histological pathology.

Mortality was increased by 13% in the 100 ppm females; moribund animals had lower spontaneous mobility, loss of coat luster, and piloerection.

The LOAEL is 100 ppm (approximately equivalent to 15 mg/kg/day) based on mortality in females. The NOAEL is 30 ppm (approximately equivalent to 4.5 mg/kg/day).

At the doses tested, there was no treatment-related increase in tumor incidence when compared to controls. Dosing was considered adequate based on decreases in survival in females.

This study is classified as acceptable/guideline and satisfies the guideline requirements (OPPTS 870.4200b; OECD 451) for a carcinogenicity study in mice.

870.5100, 870.5300, 870.5375, 870.5900 Mutagenicity

Gene Mutation

870.5100; Bacterial Gene Mutation MRID 0010440 Unacceptable/Guideline	There was no evidence of induced mutant colonies over background up to 1000 µg/plate.
870.5100; Bacterial Gene Mutation MRID 0010441 Acceptable/Guideline	There was no evidence of induced mutant colonies over background up to 5000 µg/plate.

Cytogenetics

870.5375; <i>In vitro</i> Chromosome Aberration MRID 00152692 Acceptable/Guideline	There was slight evidence of chromosome aberrations induced over background in the presence and absence of S9 activation.
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870.5385; <i>In vivo</i> Chromosome Aberration MRID 40202405 Acceptable/Guideline	There was no evidence of induced chromosome aberration over background in the presence and absence of S9 activation.
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Other Genotoxicity

870.5550; Unscheduled DNA Synthesis MRID 00152693 Acceptable/Guideline	There was no evidence that unscheduled DNA synthesis, as determined by radioactive tracer procedures, was induced.
870.5550; Unscheduled DNA Synthesis MRID 40202404 Acceptable/Guideline	There was no evidence that unscheduled DNA synthesis, as determined by radioactive tracer procedures, was induced.
870.5450; Dominant Lethal Assay MRID 00100442 Acceptable/Guideline	There was no time-related positive response of increased pre- or post-implantation loss compared to controls.
870.5915; In Vivo Sister Chromatid Exchange MRID 00152695 Acceptable/Guideline	There was a concentration-related positive response of SCE induced over background in the presence of S9-activation. A positive response of SCE induced over background was also observed in the absence of S9-activation; however, there was no clear dose-response.

Neurotoxicity

In an acute neurotoxicity study (MRID 47994201), groups of fasted 42 day-old Alpk:Ap_rSD rats 10/sex/dose were given a single oral dose of paraquat technical (33.4% w/w paraquat ion, 46.1% w/w paraquat dichloride, preparation P47) in deionized water orally (by gavage) at 10 mL/kg at doses of 0, 25, 75, or 250 mg/kg paraquat technical/kg body weight. This corresponded to doses of 0, 8.4, 25.1, and 84 mg/kg paraquat ion. Animals were observed for 14 days after dosing. Neurobehavioral assessment (functional observational battery and motor activity testing) was performed in 10/sex/group one week prior to dose administration, at approximately 2 hours after dose administration on Day 1, and at one week (Day 8) and two weeks (Day 15). At study termination, 5/sex/group were euthanized and perfused in situ for neuropathological examination. Of the perfused animals, 5/sex/group of control and 250 mg/kg animals were subjected to histopathological evaluation of brain and peripheral nervous system tissues.

No effects of the test chemical were observed in the functional observational battery, or on motor activity and nervous system histopathology.

One 250 mg/kg male dosed with paraquat technical (84 mg/kg paraquat ion) was found dead on Day 5. This male had shown a slightly reduced foot splay reflex on Days 1-4 with piloerection and “sides pinched in” on Day 4. One 250 mg/kg female was killed on Day 4, due to adverse clinical signs of irregular breathing (indicative of respiratory distress), flaccidity, “sides pinched in”, and upward spinal curvature from Days 2-4, and

piloerection and ocular discharge on Days 3-4. These deaths were considered treatment-related. All other animals survived to scheduled sacrifice. The death and respiratory distress observed in the high-dose animals are consistent with the known pulmonary toxicity of paraquat.

The LOAEL for neurotoxicity was not observed. The NOAEL is 250 mg/kg paraquat technical (84 mg/kg paraquat ion).

This neurotoxicity study is classified as **acceptable, guideline** and satisfies the guideline requirement for an acute neurotoxicity study in rats (870.6200; OECD 424).

In a subchronic neurotoxicity study (MRID 47994202) paraquat technical (33.4% (w/w) paraquat ion, 46.1% (w/w) paraquat dichloride, Batch 216, preparation reference P47) was administered to Alpk:Ap_fSD rats 12/sex/group at dose levels of 0, 15, 50, or 150 ppm (equivalent to 0, 1.0/1.1, 3.4/3.9, 10.2/11.9 mg/kg bw/day of paraquat cation in males/females) for 13 weeks. Cage side observations were recorded daily. Detailed clinical observations, including the finding of “no abnormalities detected” were recorded weekly. Neurobehavioral assessment (functional observational battery and motor activity testing) was performed in 10 animals/sex/group one week prior to dosing (pre-test) and in Weeks 1, 4, 8, and 13 of dosing. At study termination, 5/sex/group were euthanized and perfused in situ for neuropathological examination. Of the perfused animals, 5/sex/dose control and 150 ppm animals were subjected to histopathological evaluation of brain and peripheral nervous system tissues.

There were no clinical signs associated with the test material, and no effects of the test material were observed in the functional observational battery. There were also no effects of the test material on motor activity. There were also no effects of the test material on brain weights and there were no neuropathological findings.

Dosing was considered adequate, based on a previous study.

The NOAEL for subchronic neurotoxicity was 150 ppm (equivalent to 10.2-11.9 mg paraquat cation/kg in males/females). The LOAEL was not observed.

The study is classified as **acceptable, guideline** and satisfies the guideline requirement for a subchronic neurotoxicity study in rats (870.6200b).

Metabolism

870.7485 Metabolism - Rat

Non-Guideline Metabolism and Dermal Penetration

In a special study (MRIDs 00126096-00126099), a single dose of 607 µg [¹⁴C-methyl]

paraquat dichloride (99.8% radiochemical purity, Lot No. not reported) in distilled water was injected intramuscularly into each of 4 adult male Rhesus monkeys. 24-hour urine samples were collected daily for 7 days. In another experiment, a single dose of the same test compound was applied to the skin (70.0 cm²) of 6 community volunteers (ages 30-74) at approximately 11.83 µg paraquat dichloride/cm². Data were collected for bilateral applications at 3 different sites: ventral forearms, back of hands, or lower legs. The treated sites were not wrapped; the volunteers were instructed not to wash the application site for 24 hours post-treatment. Urine was collected at 4, 8, 12, and 24 hours post-dose and each consecutive 24 hours for 5 days. Urine samples were brought to the laboratory for analysis every 24 hours. In both experiments, the samples were collected in polystyrene or polypropylene containers, and 24-hour samples were stored frozen until assayed for radioactivity using liquid scintillation counting.

Monkeys eliminated 43.5-51.5% of the administered radioactivity in the urine within 24 hours post-dose and 52.3-72.3% (average 58.6%) within 7 days post-dose. Following dermal application to humans, total urinary excretion of the applied doses was 0.052-0.702% (corrected for incomplete urinary excretion with a rhesus monkey parenteral excretion factor of 58.6%). This result suggests that the compound was poorly absorbed through the skin in humans. Peak excretion occurred during the first 24 hours post-dose. Differences in absorption due to application site were not noted.

These are special studies and guidelines for the conduction of metabolism and dermal penetration studies were unavailable at the time this study was conducted. Insufficient reporting of methodology and the use of less than optimal procedures (such as not covering of the application site) suggest these data are useful for only supplementary purposes.

This study is classified as acceptable/non-guideline.

APPENDIX C. Physical/Chemical Properties

Physicochemical Properties of Technical Grade Paraquat Dichloride.		
Parameter	Value	Reference
Melting point/range	decomposes at ca. 340 °C	Product Chemistry Chapter of the Paraquat Dichloride Update, 10/10/91
pH	6.4 at 20 °C	
Density	1.5 g/cm ³ at 25 °C	
Water solubility (20 °C)	freely soluble in water: 618-620 g/L at pH 5.2, 7.2, and 9.2	
Solvent solubility (20 °C)	<0.1 g/L in acetone, dichloromethane, toluene, ethyl acetate, and hexane; 143 g/L in methanol	
Vapor pressure	<<10 ⁻⁸ kPa at 25 °C	
Octanol/water partition coefficient, Log(K _{OW})	log K _{OW} = -4.5 at 20 °C	Product Chemistry Chapter of the Paraquat Reregistration Standard, 11/25/85
Dissociation constant, pK _a	0.95 (pure active ingredient)	

APPENDIX D. Studies Reviewed for Ethical Conduct

The PHED Task Force, 1995. The Pesticide Handlers Exposure Database, Version 1.1. Task Force members Health Canada, U.S. Environmental Protection Agency, and the National Agricultural Chemicals Association, released February, 1995.

Agricultural Re-entry Task Force (ARTF) data base (SOP #3.1)

MRIDs 00126097, 00126098, & 00126099. Kelly Sherman Human Research Ethics Reviewer, OPP, June 11, 2012.